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#### Target for Antiviral Therapy

The present invention provides a crystallised module of a nuclear phosphoprotein and an assay and method for determining interactions with human papillomavirus E2 for use in drug design, for use particularly but not exclusively in designing antiviral agents with potential use in treating warts, proliferative skin lesions and carcinoma of the cervix.

#### Background to the Invention

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Human papillomaviruses (HPVs) cause warts and proliferative lesions in skin and other epithelia. In a minority of HPV types ("high risk", which include HPVs 16, 18, 31, 33, 45 and 56), further transformation of the wart lesions can produce tumours, most notably carcinoma of the cervix. HPVs have evolved a sophisticated system of control, mediated by protein:DNA and protein:protein interactions, that involves both cellular and viral proteins. The 45 kDalton nuclear phosphoprotein, E2, has two central roles in this control. It acts as the principal virally encoded transcription factor and, in association with the viral E1 protein, it creates the molecular complex at the origin of the viral DNA replication.

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E2 has three distinct modules. The N-terminal module (E2NT) of about 200 amino acids is responsible for interactions with viral and host cell transcription factors. It is followed by a flexible, proline-rich, linker module and a C-terminal module (E2CT), each of about 100 amino acids <sup>3</sup> (Fig. 1a). The E2CT binds as a homodimer to DNA sites with a consensus sequence of ACCGN<sub>4</sub>CGGT <sup>4</sup>. In most HPVs a long upstream regulatory region (URR) precedes the viral genes and contains four spatially conserved E2 binding sites: three sites proximal to the transcription start site (p97 in HPV16) and one approximately 500bp upstream.

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The dimer of E2CT serves to anchor E2 protein to its recognition sites on the DNA, the function of the E2NT is to bind and localise at least three cellular transcription



factors, Sp1, TFIIB and AMF-1, to the transcription initiation complex. In addition, E2 interacts with another viral protein, E1, which has ATPase and helicase activities. E1 itself binds to the viral origin of replication which consists of about 100 bp and is surrounded by the three E2-binding sites, proximal to the transcription start. The E2:E1 interaction greatly increases the rate of HPV genome replication<sup>2,5,6</sup>, Fig. 1a. An intact E2 is essential for the normal productive (wart) life cycle of HPV, however during malignant progression HPV DNA is integrated into the host cell genome, which usually results in disruption of the E2/E1 ORFs and loss of E2 protein, in turn leading to dysregulated expression of the viral oncogenes E6 and E7.

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Consistent with its role as a transcription regulator, E2 has been shown to direct the formation of loops in DNA containing E2 binding sites<sup>8</sup>. The loops were only formed with intact E2, and not with the E2CT alone. The E2 binding sites did not function independently and their co-operative effect was mediated by full length E2, leading the authors to suggest that there were specific interactions mediated by E2 that bridged across the set of DNA binding sites through its N-terminal. A similar DNA loop structure could also be achieved with Sp1, a cellular transcription factor, which forms a complex with distally bound E2 <sup>9</sup>; Sp1/E2 interactions are critical for transcription activation in BPV<sup>10</sup>.

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Eighty six known E2 proteins from different species and different human subtypes<sup>11</sup> are highly conserved, with sequence identities typically of 35% in the N and C-terminal modules (Fig. 1b). The crystal structure of the E2CT has been determined both alone and in complex with cognate DNA<sup>12-14</sup>. The module is a dimer with a barrel fold, and induces substantial bending (42-44°) of the DNA from its B-form double helix<sup>14</sup>.

The structure of the proteolytic fragment of HPV18 E2NT, missing 65 N-terminal residues, was recently reported at 2.1 Å spacing 15. This allowed some analysis of mutational effects on function, although the missing 65 amino acids contain residues which are essential for the transcriptional and replication activities of the protein.



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We report herein the structure of the complete E2NT determined by X-ray analysis at 1.9 Å. We have found that it is an L-shaped molecule with the residues vital for transcriptional and replication activities of the protein lying on opposite sides of the N-terminal domain. Surprisingly, our results show that the surface, vital for transcription activation, is in fact involved in association of two E2NT's into a dimer. We suggest that dimerisation of E2NT plays an important and key role in induction of DNA loop formation, the mechanism by which distally bound transcription factors would be brought close to the site of transcription initiation. More importantly, our results raise the possibility that dimer formation serves as a molecular switch between early gene expression and viral genome replication during HPV infection.

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The process of rationalised drug design requires no explanation or teaching for the skilled person but a brief description is given here of computational design for the lay reader: various computational analyses are necessary to determine whether a molecule is sufficiently similar to the target moiety or structure. Such analyses may be carried out in current software applications, such as the Molecular Similarity application of QUANTA (Molecular Simulations Inc., Waltham, Mass.) version 3.3, and as described in the accompanying User's Guide, Volume 3 pages. 134-135.

The Molecular Similarity application permits comparisons between different 20 structures, different conformations of the same structure, and different parts of the same structure. The procedure used in Molecular Similarity to compare structures is divided into four steps: 1) load the structures to be compared; 2) define the atom equivalences in these structures; 3) perform a fitting operation; and 4) analyze the 25 results.

Each structure is identified by a name. One structure is identified as the target (i.e., the fixed structure); all remaining structures are working structures (i.e., moving structures). When a rigid fitting method is used, the working structure is translated and rotated to obtain an optimum fit with the target structure. The fitting operation uses a least squares fitting algorithm that computes the optimum translation and



rotation to be applied to the moving structure, such that the root mean square difference of the fit over the specified pairs of equivalent atom is an absolute minimum. This number, given in angstroms, is reported by QUANTA.

One skilled in the art may use one of several methods to screen chemical entities or fragments for their ability to associate with a target. Again, these methods require no elucidation for the skilled person but are described here for the benefit of the unskilled reader. The screening process may begin by visual inspection of the target on the computer screen, generated from a machine-readable storage medium.

Selected fragments or chemical entities may then be positioned in a variety of orientations, or docked, within that binding pocket as defined supra. Docking may be accomplished using software such as Quanta and Sybyl, followed by energy minimization and molecular dynamics with standard molecular mechanics force fields, such as CHARMM and AMBER.

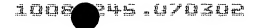
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Specialized computer programs may also assist in the process of selecting fragments or chemical entities. These include:

- GRID (P. J. Goodford, "A Computational Procedure for Determining Energetically
   Favorable Binding Sites on Biologically Important Macromolecules", J. Med. Chem.,
   pp. 849-857 (1985)). GRID is available from Oxford University, Oxford, UK.
  - 2. MCSS (A. Miranker et al., "Functionality Maps of Binding Sites: A Multiple Copy Simultaneous Search Method." Proteins: Structure, Function and Genetics, 11, pp. 29-34 (1991)). MCSS is available from Molecular Simulations, Burlington, Mass.
  - 3. AUTODOCK (D. S. Goodsell et al., "Automated Docking of Substrates to Proteins by Simulated Annealing", Proteins: Structure, Function, and Genetics, 8, pp. 195-202 (1990)). AUTODOCK is available from Scripps Research Institute, La Jolla, Calif.



4. DOCK (I. D. Kuntz et al., "A Geometric Approach to Macromolecule-Ligand Interactions", J. Mol. Biol., 161, pp. 269-288 (1982)). DOCK is available from University of California, San Francisco, Calif.

Once suitable chemical entities or fragments have been selected, they can be assembled into a single compound or complex. Assembly may be preceded by visual inspection of the relationship of the fragments to each other on the three-dimensional image displayed on a computer screen in relation to the structure coordinates of calcineurin. This would be followed by manual model building using software such as Quanta or Sybyl.

Useful programs to aid one of skill in the art in connecting the individual chemical entities or fragments include:

- CAVEAT (P. A. Bartlett et al, "CAVEAT: A Program to Facilitate the Structure Derived Design of Biologically Active Molecules". In Molecular Recognition in Chemical and Biological Problems", Special Pub., Royal Chem. Soc., 78, pp. 182-196 (1989)). CAVEAT is available from the University of California, Berkeley, Calif.
- 20 2. 3D Database systems such as MACCS-3D (MDL Information Systems, San Leandro, Calif). This area is reviewed in Y. C. Martin, "3D Database Searching in Drug Design", J. Med. Chem., 35, pp. 2145-2154 (1992).
  - 3. HOOK (available from Molecular Simulations, Burlington, Mass.).

As the skilled reader will already know, instead of proceeding to build ligand for the target in a step-wise fashion, one fragment or chemical entity at a time as described above, inhibitory or other target-binding compounds may be designed as a whole or

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de novo. These methods include:

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1. LUDI (H.-J. Bohm, "The Computer Program LUDI: A New Method for the De Novo Design of Enzyme Inhibitors", J. Comp. Aid. Molec. Design, 6, pp. 61-78 (1992)). LUDI is available from Biosym Technologies, San Diego, Calif.

- 5 2. LEGEND (Y. Nishibata et al., Tetrahedron, 47, p. 8985 (1991)). LEGEND is available from Molecular Simulations, Burlington, Mass.
  - 3. LeapFrog (available from Tripos Associates, St. Louis, Mo.).
- Other molecular modelling techniques may also be employed. See, e.g., N. C. Cohen et al., "Molecular Modeling Software and Methods for Medicinal Chemistry, J. Med. Chem., 33, pp. 883-894 (1990). See also, M. A. Navia et al., "The Use of Structural Information in Drug Design", Current Opinions in Structural Biology, 2, pp. 202-210 (1992).

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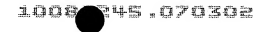
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Once a compound has been designed or selected by the above methods, the efficiency with which that entity may bind to a target may be tested and optimized by computational evaluation. For example, an effective ligand will preferably demonstrate a relatively small difference in energy between its bound and free states (i.e., a small deformation energy of binding). Thus, the most efficient ligands should preferably be designed with a deformation energy of binding of not greater than about 10 kcal/mole, preferably, not greater than 7 kcal/mole. Ligands may interact with the target in more than one conformation that is similar in overall binding energy. In those cases, the deformation energy of binding is taken to be the difference between the energy of the free entity and the average energy of the conformations observed when the inhibitor binds to the protein.

An entity designed or selected as binding to a target may be further computationally optimized so that in its bound state it would preferably lack repulsive electrostatic interaction with the target enzyme. Such non-complementary (e.g., electrostatic) interactions include repulsive charge-charge, dipole-dipole and charge-dipole



interactions. Specifically, the sum of all electrostatic interactions between the inhibitor or other ligand and the target, when the inhibitor is bound to the target, preferably make a neutral or favourable contribution to the enthalpy of binding.

Specific computer software is available in the art to evaluate compound deformation energy and electrostatic interaction. Examples of programs designed for such uses include: Gaussian 92, revision C [M. J. Frisch, Gaussian, Inc., Pittsburgh, Pa. .COPYRGT.1992]; AMBER, version 4.0 [P. A. Kollman, University of California at San Francisco, .COPYRGT.1994]; QUANTA/CHARMM [Molecular Simulations, Inc., Burlington, Mass. .COPYRGT.1994]; and Insight II/Discover (Biosysm Technologies Inc., San Diego, Calif. .COPYRGT.1994). These programs may be implemented, for instance, using a Silicon Graphics workstation, IRIS 4D/35 or IBM RISC/6000 workstation model 550. Other hardware systems and software packages will be known to those skilled in the art.

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Once the ligand has been optimally selected or designed, as described above, substitutions may then be made in some of its atoms or side groups in order to improve or modify its binding properties. Generally, initial substitutions are conservative, i.e., the replacement group will have approximately the same size, shape, hydrophobicity and charge as the original group. It should, of course, be understood that components known in the art to alter conformation should be avoided. Such substituted chemical compounds may then be analyzed for efficiency of fit to a calcineurin-like binding pocket by the same computer methods described in detail, above. Again, all these facts are familiar to the skilled person.

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Another approach is the computational screening of small molecule data bases for chemical entities or compounds that can bind in whole, or in part, to a target. In this screening, the quality of fit of such entities to the binding site may be judged either by shape complementarity or by estimated interaction energy. E. C. Meng et al., J.

30 Comp. Chem., 13, pp. 505-524 (1992).

The computational analysis and design of molecules, as well as software and computer systems therefor are described in US Patent No 5,978,740 which is included herein by reference, including specifically but not by way of limitation the computer system diagram described with reference to and illustrated in Fig 3 thereof as well as the data storage media diagram described with reference to and illustrated in Fig 4s and 5 thereof.

#### Statement of the Invention

According to a first aspect of the invention there is provided a crystallised molecular complex of an E2 N-terminal module (E2NT) dimer protein or homologue thereof, for use in rationalised drug design. We have found that the dimer comprises residues vital for transcriptional and replicational activities of said protein lying on opposite sides of an N-terminal domain, for use in rationalised drug design.

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Preferably the E2NT dimer protein is substantially as depicted in any of Figures 2c and/or 3a-d.

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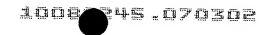
According to a second aspect of the invention there is provided an *in vitro* method for identifying and/or selecting a candidate therapeutic agent, the method comprising determining interaction of a E2 N-terminal module (E2NT) dimer in a sample by contacting said sample with said candidate therapeutic agent and measuring DNA loop formation.

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Preferably, the method is for use in identifying and/or selecting an antiviral candidate therapeutic agent.

Preferably, the candidate therapeutic agent interferes or blocks interactions of E2NT so as to interfere or block viral and/or cellular transcription factors.

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According to a third aspect of the invention there is provided use of an E2NT dimerisation inhibitor in the preparation of a medicament for use in treating warts, proliferative skin lesions and/or cervical cancer.

According to a fourth aspect of the invention there is provided a method of monitoring the efficacy of an antiviral therapy in a patient receiving a medicament for the treatment of warts, proliferative skin lesions and/or cervical cancer comprising taking a sample from said patient and measuring E2NT interactions and/or DNA loop formation.

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Thus it will be appreciated that a patient can be monitored at the start of therapy to test its effectiveness. Alternatively, a patient can be monitored once a therapy has been established so as to monitor its efficacy with a view to altering a therapy if found to be unsatisfactory.

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The human papillomavirus E2 protein controls the primary transcription and replication of the viral genome. Both activities are governed by a ~200 amino acid N-terminal module (E2NT) which is connected to a DNA binding C-terminal module by a flexible linker. The crystal structure of the E2NT module from high-risk type 16 human papillomavirus reveals an L-shaped molecule with two closely packed domains, each with a novel fold. It forms a dimer in the crystal and in solution. The dimer structure is important in the interactions of E2NT with viral and cellular transcription factors and is the key to induction of DNA loops by E2. These loops may serve to target distal DNA-binding transcription factors to the region proximal to the start of transcription. The structure has implications for antiviral drug design and cervical cancer therapy.

The invention includes method for identifying and/or selecting a candidate therapeutic agent, comprising applying rationalised drug design to a crystal structure obtainable by crystallising E2NT, cryogenically freezing the crystals and generating the crystal structure using X-ray diffraction. The method by which the E2NT crystal

structure is obtainable may comprise crystallisation using hanging-drop vapour diffusion. The method by which E2NT crystal structure is obtainable may comprise X-ray diffraction using uranium acetate and gold cyanide E2NT derivatives and refining with data extending to 1.9 Å spacing. The crystal structure may comprise the portions of amino acids Ile82, Glu90, Trp92, Lys112, Tyr138, Val145, Pro106, Lys111, Phe168, Trp134, Trp33 and Leu94. The rationalised drug design may comprise designing drugs which interact with the dimerisation surface of E2NT.

Further provided is a computer for producing a three-dimensional representation of a molecule or molecular complex, wherein said molecule or molecular complex comprises or a three-dimensional representation of a homologue of said molecule or molecular complex, wherein said homologue comprises a binding pocket that has a root mean square deviation from the backbone atoms of said amino acids of not more than 1.5Å, wherein said computer comprises:

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(a) a machine-readable data storage medium comprising a data storage material encoded with machine-readable data, wherein said data comprises the structure coordinates of E2NT amino acids Ile82, Glu90, Trp92, Lys112, Tyr138, Vall45, Pro106, Lys111, Phe168, Trp134, Trp33 and Leu94 according to Table 3;

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- (b) a working memory for storing instructions for processing said machine-readable data;
- (c) a central-processing unit coupled to said working memory and to said machinereadable data storage medium for processing said machine readable data into said three-dimensional representation; and
  - (d) a display coupled to said central-processing unit for displaying said threedimensional representation.

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In class of embodiments, the three-dimensional representation is of a molecule or molecular complex is defined by the set of structure coordinates according to Table 3, or wherein said three-dimensional representation is of a homologue of said molecule or molecular complex, said homologue having a root mean square deviation from the backbone atoms of said amino acids of not more than 1.5 Å.

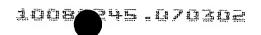
An additional aspect of the invention resides in a computer for determining at least a portion of the structure coordinates corresponding to an X-ray diffraction pattern of a molecule or molecular complex, wherein said computer comprises:

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- (a) a machine-readable data storage medium comprising a data storage material encoded with machine-readable data, wherein said data comprises at least a portion of the structural coordinates according to Table 3;
- (b) a machine-readable data storage medium comprising a data storage material encoded with machine-readable data, wherein said data comprises an X-ray diffraction pattern of said molecule or molecular complex;
- (c) a working memory for storing instructions for processing said machine-readable 20 data of (a) and (b);
  - (d) a central-processing unit coupled to said working memory and to said machinereadable data storage medium of (a) and (b) for performing a Fourier transform of the machine readable data of (a) and for processing said machine readable data of (b) into structure coordinates; and
  - (e) a display coupled to said central-processing unit for displaying said structure coordinates of said molecule or molecular complex.
- 30 A yet further aspect of the invention relates to a crystallised molecule or molecular complex comprising a dimerisation surface defined by structure coordinates of E2NT



amino acids Ile82, Glu90, Trp92, Lys112, Tyr138, Val145, Pro106, Lys111, Phe168, Trp134, Trp33 and Leu94 according to Table 3or a homologue of said molecule or molecular complex, wherein said homologue comprises a binding pocket that has a root mean square deviation from the backbone atoms of said amino acids of not more than 1.5Å. The molecule or molecular complex may be defined by the set of structure coordinates according to Table 3, or a homologue thereof, wherein said homologue has a root mean square deviation from the backbone atoms of said amino acids of not more than 1.5Å.

27. A machine-readable data storage medium (e.g. a magnetic or optical storage medium, for example a hard disc, a floppy disc or a CD-ROM), comprising a data storage material encoded with machine readable data which, when using a machine programmed with instructions for using said data, is capable of displaying a graphical three-dimensional representation of a molecule or molecular complex comprising a dimerisation surface defined by structure coordinates of E2NT amino acids Ile82, Glu90, Trp92, Lys112, Tyr138, Val145, Pro106, Lys111, Phe168, Trp134, Trp33 and Leu94 according to Table 3, or a homologue of said molecule or molecular complex, wherein said homologue comprises a binding pocket that has a root mean square deviation from the backbone atoms of said amino acids of not more than 1.5Å.

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In the machine-readable data storage medium the molecule or molecular complex may be defined by the set of structure coordinates according to Table 3, or a homologue of said molecule or molecular complex, said homologue having a root mean square deviation from the backbone atoms of said amino acids of not more than 1.5Å.

The invention further provides a machine-readable data storage medium comprising a data storage material encoded with a first set of machine readable data which, when combined with a second set of machine readable data, using a machine programmed with instructions for using said first set of data and said second set of data, can determine at least a portion of the structure coordinates corresponding to the second

set of machine readable data, wherein: said first set of data comprises a Fourier transform of at least a portion of the structural coordinates according to Table 3; and said second set of data comprises an x-ray diffraction pattern of a molecule or molecular complex.

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In another aspect, the invention resides in a method for evaluating the ability of a chemical entity to associate with a molecule or molecular complex according to the invention, comprising the steps of:

- 10 a. employing computational means to perform a fitting operation between the chemical entity and a dimerisation surface of the molecule or molecular complex; and
  - b. analysing the results of said fitting operation to quantify the association between the chemical entity and the dimerisation surface.

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# Detailed Description of the Invention

The invention will now be described by way of example only with reference to the following Figures and Tables wherein:

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Table 1 illustrates X-ray data and phasing statistics;

Table 2 illustrates refinement and model correlation;

10 Table 3 shows the structure coordinates of the E2NT module;

Figure 1a represents functional assignments of HPV 16 E2 protein;

Figure 1b illustrates sequence alignment of E2NT modules from a subset of HPV types;

Figure 2a illustrates a stereo view of electron density with a final model at the dimer interface of the E2NT module, viewed down the crystallographic two-fold axis;

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Figure 2c represents the E2NT dimer;

Figure 3a illustrates a schematic view of URR;

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Figure 3b illustrates a schematic view of loop formation induced by binding of E2 proteins to two cognate sites;

Figure 3c illustrates a model of E2 dimer formation;

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Figure 3d illustrates loops within URR as shown in Figure 3b;



Figure 4a illustrates the distribution of conserved residues on the E2NT monomer;

Figure 4b illustrates a first cluster of conserved residues on the E2NT monomer;

Figure 4c illustrates a second cluster of conserved residues on the E2NT monomer; and

Figure 4d illustrates conserved residues Gln12 and Glu39.

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Those of skill in the art understand that a set of structure coordinates for an enzyme or an enzyme-complex or a portion thereof, is a relative set of points that define a shape in three dimensions. Thus, it is possible that an entirely different set of coordinates could define a similar or identical shape. Moreover, slight variations caused by acceptable errors in the individual coordinates will have little, if any effect on overall shape. In terms of binding pockets, these acceptable variations would not be expected to alter the nature of ligands that could associate with those pockets.

The term "associating with" refers to a condition of proximity between a chemical entity or compound, or portions thereof, and a calcineurin molecule or portions thereof. The association may be non-covalent--wherein the juxtaposition is energetically favored by hydrogen bonding or van der Waals or electrostatic interactions--or it may be covalent.

The invention is also described with reference to US Patent No 5,978,740 which is included herein by reference, including specifically but not by way of limitation the computer system diagram described with reference to and illustrated in Fig 3 thereof as well as the data storage media diagram described with reference to and illustrated.

in Fig s 4 and 5 thereof.

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With reference to Figure 1a and functional assignments of E2. There is shown in a schematic view of NT, linker and CT modules of E2 indicating known functions of each module. Amino acid numbers which delimit the modules correspond to E2 from HPV16. In Figure 1b, there is shown the sequence alignment of the E2NT modules from a subset of HPV types (HPV16, HPV18, HPV11 and HPV2a) and one BPV type. Shaded blocks above the alignment indicate the experimentally determined secondary structure. Shaded blocks below the sequences indicate the minimal peptide sequences involved in protein:protein interactions, suggested by mutation studies. Residues with more than 90% identity among 86 PV types are coloured: red for internal structural residues, green for residues within the fulcrum region, blue for surface residues.

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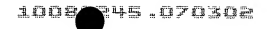
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With reference to the structural features of E2, in Figure 2a there is shown a stereo view of the electron density with the final model, at the dimer interface of the E2NT module, viewed down the crystallographic two-fold axis. The likelihood weighted map is contoured at the 1.5 σ level. Ribbons of two independent monomers are coloured blue and yellow. Side chains of ARG37 and Ile73 which are known to be critical for transactivation <sup>4,31</sup>, are shown in dark green; side chain of other residues at the dimer interface are shown in light green. Oxygen atoms are in red, nitrogen in blue, water molecules are shown as orange spheres and hydrogen bonds as dashed sticks. In Figure 2b, there is shown a stereo ribbon diagram of the E2NT module. The N1 domain is shown in aquamarine and the N2 domain in pink, with the fulcrum in green. In Figure 2c, there is shown the dimer of E2NT, showing the extent of the interface between the two subunits. The view is as in Figure 2a but rotated clockwise by 90°. Side chains of Gln12 and Glu39 which are critical for interactions with E1 <sup>31-33,37</sup> are shown in magenta. Side chains of residues at the dimer interface are coloured as per Figure 2a.

With reference to Figures 3a-d there is shown loop formation in the URR of HPV16.

30 In Figure 3a, there is shown a schematic view of the URR. The four E2-binding sites are represented by boxes. Numbers in italics indicate distances between individual



sites upstream of the p97 promoter. Two possible E2 configurations, with separate or dimeric E2NT modules are shown. In Figure 3b, there is shown a schematic view of loop formation induced by binding of E2 proteins to two cognate sites, based on the experiments reported by Knight et al<sup>8</sup>. In Figure 3d, there is shown the possible DNA loops within the URR as depicted in Figure 3b. In Figure 3c, there is shown a model of the formation of E2 dimers, showing interactions between both the C-terminal and E2NT modules. The C-terminal dimer, with its bound DNA, is based on the crystal structure of this module<sup>12</sup>. The E2NT dimer is proposed from the present work. The relative orientation and position of the E2NT and C-terminal modules is purely schematic.

With reference to Figures 4a-d there are shown functionally important residues. In Figure 4a, there is shown the distribution of conserved residues on the E2NT monomer. In Figures 4b and 4c there is shown the two clusters of conserved residues in the fulcrum of E2NT. In Figure 4d, there are shown conserved residues Gln12 and Glu39. Bonds in ball-and stick models are coloured aquamarine (N1 domain), pink (N2 domain) and green (fulcrum). Hydrogen bonds are shown as dashed lines, water molecules as orange spheres, oxygen atoms are in red, nitrogen atoms in blue and

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sulphur atoms in yellow.

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There is convincing evidence that the E2 protein has an extended structure, is flexible and that its functions depend on this property. This is probably the reason why the intact protein has not yet been crystallised in spite of intensive efforts. A major problem is the extended flexible linker module, with around 100 residues. E2NT proved difficult to crystallise, and a number of different constructs were made and overexpressed before crystallisation with residues 1 to 201 was achieved, but even this construct possessed limited stability. The protein had to be crystallised within 2-3 days of purification; crystals grew within about 48 hours but only retained useful diffraction quality for a further 2-3 days. This necessitated that crystals be rapidly vitrified in cryoprotectant buffer and stored for use as soon as detector time became available 16.

Crystals of E2NT belong to the space group P3<sub>1</sub>21 with unit-cell dimensions a=b=54.3 Å, c=155.5 Å. The structure was determined using two heavy atom derivatives and refined with data extending to 1.9 Å spacing (Fig. 2a). The main chain is well defined throughout with the exception of residues 125 and 126 which are in an exposed loop and are mobile. There was density for the last residue of the His-tag at the N-terminus, but none for the remainder of this entity. All amino acids lie in the allowed regions of the Ramachandran  $(\phi, \psi)$  plot<sup>17</sup> with 92.4% in most favoured regions<sup>18</sup>.

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The transactivation module is composed of two domains, N1 and N2, arranged so as to give it an overall L-shaped appearance. Analysis of the PDB<sup>19</sup> using DALI<sup>20</sup> shows that both have unique organisation of their secondary structures. Domain N1, which forms the N-terminus of the intact E2, is composed of residues 1 to 92, which fold into three long  $\alpha$ -helices, Figure 2 (b,c). There is a tight loop between  $\alpha 1$  and  $\alpha 2$  and a more extended one between a2 and a3. The three helices pack antiparallel to one another in the form of a twisted plane, with angles of about 20° and 25° between the pairs of consecutive helices. DALI indicated a maximum Z-score of 5.7, that could suggest a significant correlation, for colicin 1a, a membrane protein which contains three 80 Å long \alpha-helices arranged more or less coplanar<sup>21</sup>. This is the only other known protein that contains a true domain made up of such a packing of three helices. In addition there were 42 other structures which gave Z-scores above 4.0, most of which were four helix bundles, such as bacterioferritin<sup>22</sup>. However, in these only two of the three N1 helices superimposed simultaneously on two, not always adjacent, bundle helices as a result of a more planar arrangement of helices within N1. The indications are that the similarities observed reflect the optimum stacking angle of antiparallel helices against one another rather than suggesting a common ancestor for the evolution of these molecules.

Domain N2 is made up of residues 110 to 201 and is composed almost entirely of antiparallel β structure, with only one short helical segment from residues 171 to 178,



Figure 2 (b,c). The secondary structure has two short three and four stranded antiparallel  $\beta$  pleated sheets interconnected by two stranded  $\beta$  ribbons. For this domain DALI failed to identify any significant homologies to known structures, with a highest Z-score of only 2.1. From the analysis of Harris and Botchan<sup>15</sup> and the present study, the N2 fold appears to be novel.

The structure between the N1 and N2 domains (residues 93 to 109) contains two consecutive single turns of helical structure, resulting in a compact and tight turn. It packs closely against elements of both domains and is not a truly independent structural domain. Rather it forms a fulcrum in the L-shape formed by N1 and N2 where it could act as a hinge, allowing the two domains to change their relative conformation in a specific way. Several of the interactions between adjacent regions of chain in the fulcrum are mediated indirectly through H-bonds involving water molecules, suggesting the possibility of flexibility.

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One of the most striking features of the crystal structure is the association of two E2NT monomers into a tight dimer. The two E2NT monomers pack around the crystallographic 2-fold axis, as shown in Figure 2a. The dimer interface is formed mostly by amino acids from helices  $\alpha 2$  and  $\alpha 3$  of the N1 domain and by residues 142-144 from the N2 domain. The total buried surface area between the two E2NT is 2026  $A^{\circ}$ , comparable to the 2444  $A^{\circ}$  buried between the two E2CT<sup>12</sup>, which are known to form a tight dimer with a  $K_d$  of 3-6 x 10<sup>-8</sup> M <sup>23,24</sup>.

In the E2NT dimer interface, each subunit contributes a cluster of seven equivalent residues, invariant or conserved in the 86 known sequences of E2<sup>11</sup>, with many direct and water-mediated hydrogen bonds and rather few non-polar contacts, Fig. 2. Analysis of the dimer forming surfaces shows that all the direct hydrogen bonds between monomers are made through these seven amino acids. For the invariant Arg37, all possible side-chain hydrogen bonds are made and all are well defined, Figure 2. Three of them are across the dimer interface. One hydrogen bond is critical, from NH2 to the main chain carbonyl oxygen of Leu77. A second hydrogen bond from NH2 is to OG1 of Thr81; in five out of 86 sequences this residue is



glutamine, and modelling shows a hydrogen bond is possible to the NE of Arg37. The NH1 of Arg71 H-bonds to the OE1 of residue 80, which is Glu or Gln in all but six variants. At the NE of Arg37 there is an ideal H-bond to water that itself makes another strong H-bond across the dimer interface to the main-chain carbonyl oxygen of residue 142. The role of the invariant Ile73 is the filling of the intersubunit nonpolar volume made up of the aliphatic parts of Arg37, Gln76 and of Leu77 - in this case from both monomers. The Leu77 is in a few sequences substituted by valine or isoleucine and in 9 out of 86 known sequences by methionine. Inspection of the structure shows that Leu77 is partially exposed to the solvent and therefore different hydrophobic side chains could be easily accommodated at this site. important non-polar side chain is Ala69. Its side chain methyl packs into the surface of the other monomer at van de Waals distance from the main chain of residue 142. The only observed mutation of Ala69 is to Gly, and is easily accommodated. Gln76 is conserved or has homologous substitutions in about 2/3 of E2 sequences; in about 1/4 of the sequences there is methionine or valine at this position<sup>11</sup>. Although hydrophobic substitutions of Gln76 would disrupt the hydrogen bonding to Glu80 across the dimer interface, and to Arg37 from the same subunit, the hydrophobic side chain at residue 76 could instead make a compensating hydrophobic interaction with the adjacent intersubunit hydrophobic pocket formed by Ile73 and Leu77.

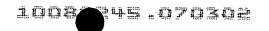
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Modelling of the amino acid variations in the 86 known papillomavirus E2 proteins into the other contacts at the dimer interface shows that they generally can be accommodated (data not shown). The consistency of the hydrogen bonds and van de Waals contacts at the monomer-monomer interface in the various sequences suggests therefore that the E2NT dimer interactions are potentially present in all papillomaviruses.

The first experimental evidence for the E2NT dimerisation in the presence of DNA with multiple E2-binding sites was provided by Knight et al in 1991<sup>8</sup>. Their studies showed that intact E2 led to the formation of DNA loops on templates with widely separated E2 binding sites, while a truncated E2, containing the DNA-binding E2CT but missing the N-terminal 161 residues, did not. Such dimerisation is further



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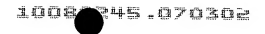
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supported by the observed synergistic transcription activation by a complex of two DNA-bound E2 dimers<sup>25</sup>.

To analyse the functional behaviour of the E2NT dimers further, we measured the constant bу sedimentation equilibrium ultracentrifugation of recombinant E2NT protein containing the 201 N-terminal amino acids. A value of  $K_d = 8.1 \pm 4 \times 10^{-6}$  M was obtained, indicating mediumstrength association. The micromolar range of the E2NT dimer  $K_d$  is certainly physiologically significant, and compares well with values for other transcription factors which have relatively low dissociation constants, often with the  $K_d$  values between 1 µM and 20 µM <sup>26,27</sup>. In vivo, the interaction could be enhanced when the two E2NT modules are placed in close proximity. Indeed, E2CT forms dimers which bind to the multiple DNA-binding sites located within the URR of viral DNA with  $K_d$  of protein:DNA interactions usually in the nanomolar range<sup>28</sup>. Consequently, the local concentration of E2NT, bound to the E2CT via the non-conserved, flexible ~80 amino-acid linker, is effectively increased.

E2NT dimer interactions, as seen in the crystal structure, could form either between modules which are already part of a single E2 dimer, formed as a result of E2CT dimerisation interactions and bound to a single E2 binding site on the DNA (Fig. 3a), or between two preformed E2 dimers located on different E2 binding sites (Fig. 3b). The results of the electron microscopy suggest that the latter dimerisation does occur<sup>8</sup>. Although no direct experimental evidence exists for the former dimerisation, it does also seem possible due to the flexibility of the linker connecting the two modules. We propose that E2 molecules may initially keep their N-terminal modules within their internal dimers, but swap N-terminal modules and cross link to E2 molecules bound to distant DNA binding sites to form active loop structures during transcriptional activation and / or HPV DNA replication (Figure 3d). As discussed below, the effects of mutations on transcriptional transactivation can be explained in terms of the dimer being an essential element in this process.



E2 is a regulator of both transcription and viral DNA replication and thus interacts with other viral and host macromolecules in the infected cell. Indication of the possible importance of individual residues in the function comes firstly from the structure, secondly from the extensive set of sequences of the papillomaviral E2's and thirdly from mutagenesis studies on the individual proteins. In the following we make a primary attempt to map the molecule's function onto its structure.

The pattern of amino acid conservation for the 86 available papilloma sequences<sup>11</sup> has been analysed using the GCG program suite<sup>29</sup>. The sequences exhibit striking variation, characteristic of some virus families. However, 33 of the total 201 residues in the E2NT construct were totally or highly conserved. Fig. 4a illustrates the distribution of these 33 residues in the dimer. These were categorised into two sets: those with an essentially structural role and those exposed on the surface with a potential for intermolecular interactions. Thirteen residues (Fig. 1b) are buried or play a purely structural role within the monomer, they are not expected to be of functional importance and will not be discussed here.

A further 12 of these 33 residues stand out as having a structural role in the interface of the N1 and N2 domains. They form three clusters, the first making direct interactions between the two domains (Ile82, Glu90, Trp92, Lys112, Tyr138, Val145) and two separate sets of interactions, one from N2 (Pro106, Lys111, Phe168, Trp134) and the other from N1 (Trp33, Leu94) to the structure connecting them, referred to here as a fulcrum. The first two clusters are shown in Figure 4 b, c and it can be seen that Lys111 and Lys112 play key roles. Their side chains point in opposite directions to one another and their terminal amino groups are involved in near ideal patterns of hydrogen bonds. The flat surfaces of their extended side chains stack against Trp134 and Trp92, respectively. This clustering of invariant residues at the interface indicates a functional importance for the relative orientation of N1 and N2. The fulcrum could indeed provide a flexible pivot between the two domains, but there is no direct evidence for this as yet. Finally, while the side chain of Glu90 is held tightly in place by two H-bonds and could have a structural role, its OE2 atom is



exposed on the surface and is surrounded by near invariant side-chains, which may thus play a part in interactions with other molecules.

Of the remaining eight conserved residues, mutational substitutions of Glu20, Glu100 and Asp122 <sup>30-33</sup> had moderate effects on the transactivation and replication properties of E2, which depended on a particular viral strain. Glu20 lies on the top surface of N1. Asp122 lies far away on the distal surface of N2. Glu100 is completely exposed and points into the solvent at the junction of the L between the N1 and N2 domains. The functional role of these amino acids has yet to be clarified.

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Three conserved amino acids (Arg37, Glu39 and Ile73) have been subjected to point mutation and the effects on the two principal functions of E2, i.e. transactivation and HPV DNA replication have been assessed (reviewed in<sup>4</sup>,also <sup>31,34,35</sup>). Together with the remaining two conserved amino acids, Gln12 and Ala69, these residues form two functionally important surfaces (see below).

Finally, a number of the mutational results (reviewed in <sup>4</sup>, also <sup>31,34,35</sup>) correspond to residues that can be assigned to structural roles. Substitution of these residues will lead to substantial conformational changes and a probable inability to fold correctly. This is particularly true for some of the deletion mutants involving the core of the molecule. Knowledge of the structure will allow a more rational choice and design

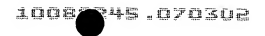
of mutants in the future.

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The induction of DNA loops by E2NT dimerisation could be important for the construction of the active transcription bubble by targeting DNA-binding transcription factors, bound at distal sites, to the region proximal to the start of transcription (reviewed in <sup>36</sup>). In support of this, residues Arg37, Ile73 and Gln76 map onto the surface of E2NT involved in dimer formation, and mutations result in considerable disruption of transactivation, while having little effect on replication, <sup>4,15,31</sup>. The structure also shows that Ala69 which points its side chain methyl across the dimer interface, is also critical for transactivation. Mutational substitutions to



amino acids with longer side chains should have a knock out effect on E2NT dimer formation and consequently on transactivation.

The sites of association with cellular transcription factors AMF-1 (residues 74-134) and TFIIB (134-216) were previously mapped onto the E2NT module (Figure 1) using a series of deletion mutants as well as point mutations<sup>34,35</sup>. These sites were mutually exclusive. In the structure, residues 74-134 include the fulcrum, while residues 134-216 correspond to domain N2. Further biochemical and structural studies can now be planned to characterise these interactions in more detail.

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Replication of the viral genome is initiated by binding of another viral protein, E1, to the origin of DNA replication<sup>4</sup> which is itself flanked by two E2 binding sites, Fig. 3a. While the function of E2CT dimers is to bind specifically to the DNA sites, E2NT interaction with E1 enhances the binding of E1 to this region. Mutational substitutions of Glu39 generally retained transcriptional activation while DNA replication was substantially reduced<sup>31-33,37</sup>. In the structure, the conserved Glu39 makes every possible hydrogen bond by its side chain carboxyl oxygens, Fig. 4d. One hydrogen bond is to NE2 of Gln12, which is absolutely conserved in all known sequences of E2. The other three hydrogen bonds are to the water molecules which are part of an intimate net of well-defined water molecules surrounding Glu39 and mediating its interactions with adjacent residues. Interestingly, a number of these protein interactions with water molecules are conserved as they are made to the protein backbone, including carbonyl oxygens of Gln12, Met36 and Lys68. While mutation of Gln12 in BPV1 only slightly affected both transactivation and replication, it substantially reduced cooperative origin binding<sup>30,32</sup>. The close positioning of Gln12 and Glu39 in the three-dimensional structure further enhances the notion that these two resides are involved in interactions with E1. The conserved set of interactions at Gln12/Glu39 suggests that the main chain carbonyl oxygens of Gln12 and Met36 and the conserved water molecules could be also involved in these interactions. Gln12/Glu39 are surrounded by Leu8, Ile15, Met36, Tyr43, Gln57 and



Lys68, which are unlikely to contribute into E2/E1 interactions, as these residues are not well conserved in E2 sequences from different papillomaviruses.

The Gln12/Glu39 cluster lies on a side of the N1 domain which is opposite to the side involved in transactivation (and dimerisation), Figure 2c. Notably, the spatial separation of the two functionally important surfaces suggests that E2NT module could be able to interact with E1 at the same time as it interacts through the dimerisation interface with another E2NT module.

10 The structure reported here for the entire E2 transactivation module, has several implications for understanding of E2 function. It is now possible to map known mutations onto the E2 three-dimensional structure, and to use the knowledge of amino acid conservation and the effects of mutations to assign roles in folding, structure and function to residues. To this end, our results indicate that molecular surfaces involved in transactivation and E1-binding are located at opposite sides of the N1 domain of E2NT, suggesting that both surfaces could be accessed simultaneously by other protein factors. In line with these observations, E1 has been shown to modulate transactivation by directly interacting with E2, leading to repression of transactivation in the presence of excess E1<sup>38</sup>. It is not inconceivable that the docking of E2NT dimer with E1 is sufficient to block further association with other target proteins.

The structure shows that the transactivation surface is involved in the formation of the E2NT dimer, which could cross-link E2 molecules bound by their E2CT modules to well-separated DNA sites. Inevitably, such dimerisation would cause DNA to form a loop structure, targeting distally bound transcription factors to regions close to the promoter. While this process has been suggested to be essential for transactivation<sup>36</sup>, the definition of interacting surfaces between E2 and other cellular transcription factors requires a great deal of further study.

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Our results suggest that the process of DNA loop formation could involve swapping of E2NT modules across E2 dimers bound at separated DNA sites (Fig. 3a-d). The polar components of the monomer-monomer interactions may favour such exchange. Domain swapping is a well-recognised phenomenon that occurs relatively frequently between two individual monomers containing domains connected by a flexible linker <sup>39,40</sup>. E2 is to our knowledge the first example where the swapping event is predicted to occur between dimers.

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The dimerisation surface of E2 represents a good target for designing anti-viral drugs, since it is essential for viral transcription, there is no homologous human protein and the residues forming the interface are highly conserved among different viral strains. Dynamic interactions between transcription factors play a central role in the regulation of transcription and replication. Dimerisation, heterodimerisation and the monomer-to-dimer transition may play important roles during the control of the papillomavirus life cycle. These processes themselves can be regulated through phosphorylation, proteolysis, interaction with small ligands or changes in their intracellular concentration. It has been suggested that E2 can regulate the switch between early gene expression and viral genome replication during HPV infection<sup>41</sup>. It is possible that dimerisation of E2NT modules plays an essential role during this process. One scenario would be to activate transcription via induction of DNA loop formation at early stages of the viral life cycle. At later stages, when the concentration of expressed E2 proteins within the cell becomes high and comparable with the Kd for E2 dimer formation, free E2NT modules could compete for dimerisation with those involved in DNA loop formation and titrate them away, switching off transcription and stimulating replication. It is also possible that other protein factors could be involved in this process, including, for example, E1.

The invention therefore includes the use of E2NT crystal structure in the design of anti-viral drugs, since it is essential for viral transcription. In the rationalised computational design of drugs using the crystal structure, computational analyses are therefore necessary to determine whether a molecule or the E2NT-binding portion



thereof is sufficiently similar to the E2NT structure. Such analyses may be carried out in current software applications, such as the Molecular Similarity application of QUANTA (Molecular Simulations Inc., Waltham, Mass.) version 3.3, and as described in the accompanying User's Guide, Volume 3 pages. 134-135.

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The Molecular Similarity application permits comparisons between different structures, different conformations of the same structure, and different parts of the same structure. The procedure used in Molecular Similarity to compare structures is divided into four steps: 1) load the structures to be compared; 2) define the atom equivalences in these structures; 3) perform a fitting operation; and 4) analyze the results.

Each structure is identified by a name. One structure is identified as the target (i.e., the fixed structure); all remaining structures are working structures (i.e., moving structures). Atom equivalency within QUANTA is defined by user input and, for the purpose of this invention equivalent atoms may be defined as protein backbone atoms (N, C.alpha., C and O) for all conserved residues between the two structures being compared. We will also consider only rigid fitting operations.

When a rigid fitting method is used, the working structure is translated and rotated to obtain an optimum fit with the target structure. The fitting operation uses a least squares fitting algorithm that computes the optimum translation and rotation to be applied to the moving structure, such that the root mean square difference of the fit over the specified pairs of equivalent atom is an absolute minimum. This number, given in angstroms, is reported by QUANTA.

For the purpose of one class of embodiments this invention, any set of structure coordinates of a molecule or molecular complex that has a root mean square deviation of conserved residue backbone atoms (N, C.alpha., C, O) of less than 1.5 .ANG. when superimposed—using backbone atoms—on the relevant structure coordinates of E2NT are considered identical. More preferably, the root mean square



deviation is less than 1.0 .ANG.. Most preferably, the root mean square deviation is less than 0.5 .ANG..

The term "root mean square deviation" means the square root of the arithmetic mean of the squares of the deviations from the mean. It is a way to express the deviation or variation from a trend or object. For purposes of this invention, the "root mean square deviation" defines the variation in the backbone of a protein from the backbone of E2NT a dimerising portion thereof, for example as defined by the structure coordinates of E2NT described herein.

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The term "least squares" refers to a method based on the principle that the best estimate of a value is that in which the sum of the squares of the deviations of observed values is a minimum.

#### 15 Materials and Methods

#### Purification and crystallisation.

Details of the purification and crystallisation of E2NT have been described previously <sup>16</sup>. Briefly, the ORF encoding the N-terminal 201 residues of HPV-16 E2 was cloned into the prokaryotic expression plasmid pET15b downstream of the 20-residue His-tag leader sequence; protein was expressed in *E. coli*BL21(DE3)pLysS and purified using nickel affinity and anion exchange chromatography. Crystals were obtained by hanging drop vapour diffusion with 0.8-1.2M ammonium sulphate, 0.1M triethanolamine pH 8.0-8.3 and 3-5% 2-methyl-2,4-pentanediol. Crystals grew only with very fresh protein preparations and deteriorated in terms of diffraction quality in less than a week. This necessitated freezing and storage of crystals in liquid nitrogen immediately after growth, as discussed above.

#### Structure determination.

All data were recorded on cryogenically frozen crystals. A native crystal was frozen for which initial data were recorded to 3.4 Å<sup>16</sup>. For the screening of derivatives,

crystal stability was even more limiting. Nine crystals were soaked in various heavy atom reagents immediately after growth. The crystals were screened in-house using a MAR research imaging plate on a Rigaku RU200 rotating anode source, by recording 3° of data for each and analysing the fractional isomorphous difference from the native. Three derivatives showed promising differences from the native, in the range of 15-20% after scaling using SCALEPACK<sup>42</sup> and were stored in liquid nitrogen. The native crystal was transported to EMBL Hamburg where 1.9 Å data were measured using synchrotron radiation from beam line X11, Table 1. In addition data were recorded at EMBL for the three promising derivatives to about 2.7 Å. Two of these derivatives proved useful in phase determination and the structure was solved by multiple isomorphous replacement with anomalous scattering (MIRAS) at 2.7 Å. The two derivatives were solved independently using the CCP4 suite 43 from the difference Patterson synthesis and by direct methods as implemented in SHELX<sup>44</sup>. Both contained a single heavy atom site. Phases, calculated using MLPHARE, were enhanced by solvent flattening4<sup>5</sup> using a solvent content of 50 %. The resulting high quality density map was easily interpretable and the initial model was built using QUANTA (Molecular Simulations) for all but four residues of the construct, ignoring the His-tag. The model was completed with REFMAC (resolution 20-1.9 Å) using a bulk solvent correction, to an R-factor of 23.3 % (R<sub>free</sub> 29.7 % - for 5 % of the data). There are 221 residues in the recombinant protein: the first twenty comprise the His-Tag. The final model contains all but two of the 201 residues of the real protein: residues 125-126 are disordered and lie in a flexible surface loop. Only one residue, His0, of the His-tag has clear density and an ordered conformation. In addition there are 187 water molecules, which were selected using ARP<sup>46</sup>during the course of refinement. The main statistics of the refined model are shown in Table 2.

### Analytical ultracentrifugation.

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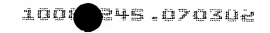
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Experiments were carried out in an Optima XL-A ultracentrifuge (Beckman-Coultier, CA, USA) using scanning UV optics. During the experiments, the recombinant E2NT was in 10mM TrisHCl pH 8.0, 5mM DTT, 0.2 mM EDTA, 300 mM NaCl.



Data were obtained at rotor speeds of 12,000 and 16,000 rpm, and the time to equilibrium was 10-12 hours. All runs were carried out at 293K, and all radial scans were at a wavelength of 280 nm. Dissociation constants were obtained by nonlinear regression using the Beckman ultracentrifuge software.

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Table 1
X-ray data and phasing statistics

Data set	Native	UAc	AuCN			
Space Group	P3 <sub>1</sub> 21	P3 <sub>1</sub> 21	P3 <sub>1</sub> 21			
	54.68	54.49	54.58			
a ,b (Å)		155.66	156.50			
c (Å)	155.73		20 - 2.7			
Resolution (Å)	30-1.9	20-2.7				
Temperature, K	120	120	120			
Wavelength (Å)	0.86	0.86	7937			
Unique reflections	21751	7873				
Completeness (%)	98.8 (89.3)	99.8 (96.1)	99.7 (93.8)			
(outer shell)			2 2 (1 (0 2 (0))			
R-merge (outer shell)	0.058 (0.339)	0.073 (0.271)	0.061 (0.268)			
Phasing Power: (centri	c / acentric)	1.55 / 2.07	0.95 / 1.40			
FOM: MIRAS		0.59				
FOM: DM 20-2.7 Å (2	2.7 - 1.9 Å)	0.88 (0.61)	0.88 (0.61)			
DM: Mean phase chan		32 °	32 °			
R-factor (FreeR)	0.223 (0.295)		·			

# Table 2

### Refinement and model correlation

	Resolution		1.9 - 10.0 Å							
	Number of protein atoms		1622							
5	Number of solvent sites		211							
	Number of reflections used in refinement		20637							
	Number of reflections used for Rfree calculation	1111								
	R-factor ‡		0.232							
	Rfree ‡		0.305							
10	Average atomic B-factor*, Å <sup>2</sup>	protein atoms	38.0							
		water molecules	48.5							
	R.m.s. deviations from ideal geometry (Å). Targets in parentheses									
		bond distance	0.013 (0.020)							
		angle distance	0.026 (0.040)							
15		chiral volume	0.142 (0.200)							

 $<sup>\</sup>label{eq:crystallographic R-factor, R_free} \begin{tabular}{l} $\updownarrow$ Crystallographic R-factor, $R_{free}$ = $\sum \|F_o\| - |F_c\| / \sum |F_o| \;. \end{tabular}$ 

20 Table 3

	CRYST	54.	680	54	. 680	155.73	0 90.00 90.00	) 120.00 P31	21
	SCALE1		0.0	1829	0.	.01056	0.00000	0.00000	
25	SCALE2		0.0	0000	0.	.02112	0.00000	0.00000	
	SCALE3		0.0	0000	0.	. 00000	0.00642	0.00000	
	ATOM	1	N	HIS	A	0	5.469 -26.512	52.262 1.	00 61.92
	ATOM	2	CA	HIS	Α	0	6.434 -25.669	51.568 1.	00 61.84
	ATOM	3	С	HIS	A	0	6.263 -25.743	50.051 1.	00 53.91
30	ATOM	4	Ó	HIS	A	0	6.089 -24.713	49.607 1.	00 69.59
	ATOM	5	CB	HIS	Α	0	7.837 -26.127	51.965 1.	00 54.18
	ATOM	6	CG	HIS	A	0	7.848 -26.468	53.431 0.	00 99.00
	ATOM	7	ND1	HIS	Α	0	7.914 -25.533	54.412 0.	00 99.00
	ATOM	8	CD2	HIS	A	0	7.732 -27.728	54.027 0.	00 99.00
35	ATOM	9	CE1	HIS	A	0	7.828 -26.215	55.570 0.	00 99.00
	ATOM	10	NE2	HIS	A	0	7.723 -27.531	55.370 0.	00 99.00
	MOTA	11	N	MET		1	6.663 -26.896	49.478 1.	00 56.24
	ATOM	12	CA	MET	A	1	6.435 -27.076	48.053 1.	00 56.42
	ATOM	13	C	MET		1	5.209 -26.282	47.619 1.	00 56.07
40	ATOM	14	ō	MET		1	5.293 -25.299	46.911 1.	00 56.51
	ATOM	15	СВ	MET		ī	6.216 -28.565	47.788 1.	00 60.46
	ATOM	16	CG	MET	-	1	6.856 -29.020	46,477 0.	00 99.00
	ATOM	17	SD	MET	-	ī	7.244 -30.775	46.483 0.	00 99.00
	ATOM	18	CE	MET		ī	7.499 -30.975		00 99.00
45	ATOM	19	N	GLU		2	4.035 -26.755		00 54.92
	ATOM	20	CA	GLU		2	2.803 -26.044	47.744 1.	00 53.59

				MEM 1			6.856 -29.02	0 46.477	0.00 99.00
	ATOM	16	CG	MET A		1			0.00 99.00
	ATOM	17	SD	MET A		1	7.244 -30.77	-	
	MOTA	18	CE	MET A		1	7.499 -30.97		0.00 99.00
	ATOM	19	N	GLU A		2	4.035 -26.75		1.00 54.92
5	MOTA	20	CA	GLU A		2	2.803 -26.04		1.00 53.59
	ATOM	21	С	GLU A	A 2	2	2.870 -24.57		1.00 52.81
	MOTA	22	0	GLU A	A :	2	2.555 -23.66		1.00 51.69
	MOTA	23	CB	GLU Z	A :	2	1.661 -26.74	0 48.482	1.00 56.88
	ATOM	24	CG	GLU Z		2	2.090 -28.09	2 49.054	0.00 99.00
10	ATOM	25	CD	GLU A		2	1.019 -28.61	.0 49.983	0.00 99.00
	ATOM	26		GLU Z		2	0.454 -27.81	9 50.722	0.00 99.00
	ATOM	27	OE2		A :	2	0.761 -29.81	1 49.963	0.00 99.00
	ATOM	28	N	THR		3	3.260 -24.34		1.00 52.06
	ATOM	29	CA	THR A		3	3.300 -22.98		1.00 51.61
15	ATOM	30	C	THR A	Α .	3	4.161 -22.05		1.00 50.30
13	ATOM	31	ŏ	THR	Δ .	3	3.731 -21.00		1.00 49.91
	ATOM	32	CB	THR A		3	3.858 -23.02		1.00 54.31
				THR A		3	2,975 ~23.78		1.00 56.98
	ATOM	33				3	3.960 -21.60		1.00 55.18
20	ATOM	34		THR A		4	5.372 ~22.49		1.00 50.11
20	ATOM	35	N	LEU A			6.201 -21.69		1.00 50.48
	MOTA	36	CA	LEU A		4	5.553 -21.53		1.00 50.40
		. 37	C	LEU .		4 .			1.00 50.73
	ATOM	38	0	LEU A		4	5.520 -20.41		
	ATOM	39	CB	LEU I		4	7.603 -22.28		1.00 52.72
25	ATOM	40	CG	LEU A		4	8.545 -22.25		1.00 56.58
	ATOM	41		LEU A		4	9.819 -23.03		1.00 55.37
	MOTA	42	CD2	LEU A		4	8.829 -20.82		1.00 56.27
	MOTA	43	N	CYS Z		5	5.028 -22.61		1.00 49.77
	ATOM	44	CA	CYS 7		5	4.362 -22.53		1.00 48.93
30	ATOM	45	С	CYS 2	Α :	5	3.218 -21.53	44.589	1.00 48.27
	ATOM	46	0	CYS Z	A !	5 .	3.136 -20.69	8 43.682	1.00 47.03
	ATOM	47	CB	CYS Z	A	5	3.865 -23.87	9 44.075	1.00 49.50
	ATOM	48	SG	CYS Z	A. !	5	5.217 -24.97	2 43.626	1.00 50.79
	ATOM	49	N	GLN A		6	2.356 -21.62	7 45.610	1.00 46.85
35	ATOM	50	CA	GLN A		6	1.227 -20.73	8 45.675	1.00 47.22
	ATOM	51	C	GLN A		6	1.666 -19.27	6 45.865	1.00 46.83
	ATOM	52	Õ	GLN A		6	1.050 -18.38		1.00 48.60
	ATOM	53	СB	GLN A		6	0.272 -21.07		1.00 50.54
	ATOM	54	CG	GLN A		6	-0.681 -22.22		1.00 55.34
40	ATOM	55	CD	GLN A		6	-1.144 -22.87		1.00 59.63
40	ATOM	56		GLN A		6	-1.101 -22.22	2 48.853	1.00 61.52
				GLN A		6	-1.482 -24.15		1.00 57.45
	ATOM	57				7	2.628 -19.03		1.00 46.33
	ATOM	58	N	ARG I		7	3.162 -17.70		1.00 46.38
45	ATOM	59	CA	ARG I		7	3.780 -17.14		1.00 44.90
43	ATOM	60	C	ARG A		7	3.544 -15.98		1.00 45.29
	ATOM	61	0	ARG A			4.267 -17.68		1.00 50.84
	ATOM	62	CB	ARG		7			1.00 62.38
	ATOM	63	CG	ARG A		7	3.690 -17.86		1.00 02.30
	ATOM	64	CD	ARG A		7	2.884 -16.61		
50	ATOM	65	NE	ARG A		7	3.786 -15.53		
	MOTA	66	CZ	ARG A		7	3.406 -14.30		1.00 83.50
	MOTA	67		ARG I		7	2.122 -13.96		1.00 85.24
	ATOM	68	NH2	ARG A		7	4.316 -13.41		1.00 85.59
	ATOM	69	N	LEU A		8	4.608 -17.91		1.00 45.22
55	MOTA	70	CA	LEU A		В	5.212 -17.42		1.00 45.85
	ATOM	71	С	LEU 2	A (	В	4.161 -17.13		1.00 46.67
	ATOM	72	0	LEU Z		В	4.197 -16.05	5 42.025	1.00 46.25
	ATOM	73	CB	LEU Z		В	6.185 -18.47	7 43.194	r.00 41.52
	ATOM	74	CG	LEU 2		В	6.979 -18.05		1.00 44.17
60	ATOM	75		LEU I		В	7.941 -16.92		1.00 44.96
	ATOM	76		LEU Z		8	7.723 -19.24		1.00 44.18
	ATOM	77	N	ASN Z		9	3.193 -18.01	4 42.508	1.00 47.73
	ATOM	78	CA	ASN Z		9	2.065 -17.82	5 41.610	1.00 48.73
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	MOTA	79	С	ASN A	9		. 351	-16.516	41.925	1.00 48.98
	MOTA	80	ō	ASN A	9			-15.727	40.999	1.00 48.98
	ATOM	81	СВ	ASN A	9			-18.923	41.725	1.00 54.35
	ATOM	82	CG	ASN A	وَ			-20.167	40.912	1.00 58.01
5		83		ASN A	· 9			-20.427	40.356	1.00 60.06
2	MOTA	84		ASN A	9			-20.991	40.841	1.00 63.02
	ATOM			VAL A	10			-16.267	43.206	1.00 48.31
	ATOM	85	N	VAL A	10	-		-15.003	43.528	1.00 48.22
		86	CA	VAL A	10			-13.837	43.252	1.00 48.34
10	MOTA	87 88	0	VAL A	10			-12.816	42.688	1.00 47.81
10	ATOM	89	СВ	VAL A	10			-14.918	44.981	1.00 53.07
	ATOM	90		VAL A	10			-13.487	45.353	1.00 53.77
	ATOM ATOM	91		VAL A	10			-15.827	45.176	1.00 54.90
	ATOM	92	N	CYS A	11			-14.011	43.661	1.00 47.68
15	ATOM	93	CA	CYS A	11			-12.938	43.426	1.00 47.78
13	ATOM	94	C	CYS A	11			-12.615	41.954	1.00 47.29
	ATOM	95	ŏ	CYS A	11			-11.473	41.499	1.00 47.31
	ATOM	96	СВ	CYS A	11			-13.269	44.144	1.00 48.13
•	ATOM	97	SG	CYS A	11			-11.884	44.082	1.00 44.06
20	ATOM	98	N	GLN A	12			-13.633	41.120	1.00 47.63
20	ATOM	99	CA	GLN A	12			-13.484	39.702	1.00 48.32
	ATOM	100	C	GLN A	12			-12.946	38.951	1.00 48.82
	ATOM	101	Õ	GLN A	12			-12.258	37.946	1.00 48.64
	ATOM	102	CB	GLN A	12			-14.783	39.092	1.00 45.97
25	ATOM	103	CG	GLN A	12			-15.213	39.590	1.00 45.90
23	ATOM	104	CD	GLN A	12			-16.359	38.862	1.00 46.71
	ATOM	105		GLN A	12			~17.320	38.425	1.00 45.33
	MOTA	106		GLN A	12			-16.294	38.702	1.00 49.43
	ATOM	107	N	ASP A	13			-13.199	39.470	1.00 48.93
30	ATOM	108	CA	ASP A	13		0.516	-12.691	38.853	1.00 49.49
	ATOM	109	C	ASP A	13			-11.198	39.085	1.00 49.55
	ATOM	110	ō	ASP A	13		0.082	-10.444	38.171	1.00 49.73
	ATOM	111	CB	ASP A	13	-1	0.732	-13.392	39.411	1.00 52.95
	ATOM	112	CG	ASP A	13	-1	0.955	-14.680	38.932	0.00 99.00
35	ATOM	113		ASP A	13	-1	0.110	-15.160	38.175	0.00 99.00
	ATOM	114		ASP A	13			-15.191	39.132	0.00 99.00
	ATOM	115	N	LYS A	14	(	0.801	-10.735	40.269	1.00 48.95
	ATOM	116	CA	LYS A	14		0.809	-9.313	40.556	1.00 49.25
	ATOM	117	C	LYS A	14		1.794	-8.575	39.658	1.00 49.07
40	ATOM	118	0	LYS A	14		1.470	-7.519	39.119	1.00 49.78
-	ATOM	119	CB	LYS A	14		1.109	-9.040	42.030	1.00 52.53
	ATOM	120	CG	LYS A	14		0.070	-8.421	42.768	1.00 62.87
	ATOM	121	CD	LYS A	14		0.269	-6.975	42.329	1.00 66.42
	ATOM	122	CE	LYS A	14		1.227		43.257	1.00 70.58
45	ATOM	123	NZ	LYS A	14		0.835		43.452	1.00 72.01
	ATOM	124	N	ILE A	15	:	2.984		39.468	1.00 48.82
	ATOM	125	CA	ILE A	15		3.992		38.595	1.00 49.56
	ATOM	126	С	ILE A	15		3.467		37.165	1.00 50.17
	MOTA	127	0	ILE A	15		3.538		36.561	1.00 50.19
50	ATOM	128	CB	ILE A	15	ļ	5.288		38.669	1.00 43.08
	MOTA	129	CG1	ILE A	15		5.931		40.054	1.00 45.94
	MOTA	130	CG2	ILE A	15		6.286		37.597	1.00 47.46
	MOTA	131	CD1	ILE A	15			-10.120	40.472	1.00 42.94
	ATOM	132	N	LEU A	16		2.880		36.623	1.00 51.16
55	ATOM	133	CA	LEU A	16		2.272		35.291	1.00 52.54
	ATOM	134	С	LEU A	16		1.135		35.191	1.00 52.81
	MOTA	135	0	LEU A	16		1.023		34.194	1.00 53.37
	MOTA	136	CB	LEU A	16			-10.810	34.847	1.00 56.20
	MOTA	137	CG	LEU A	16			-11.696	34.504	1.00 61.93
60	MOTA	138		LEU A	16			-13.139	34.904	1.00 65.17
	MOTA	139		LEU A	16			-11.572	33.041	1.00 62.31
	ATOM	140	N .	THR A	17		0.274		36.204	1.00 52.95
	ATOM	141	CA	THR A	17	-(	0.789	-7.332	36.217	1.00 53.67

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	MOTA	142	С	THR A	17		-0.232	-5.916	36.173	1.00 54.28
	ATOM	143.	0	THR A	17		-0.860	-5.026	35.590	1.00 54.21
	ATOM	144	CB	THR A	17		-1.677	-7.468	37.468	1.00 55.51
	ATOM	145	OG1	THR A	17		-2.321	-8.742	37.469	1.00 54.73
5	ATOM	146	CG2	THR A	17		-2.713	-6.355	37.551	1.00 52.35
•	ATOM	147	N	HIS A	18		0.879	-5.647	36.878	1.00 53.99
	ATOM	148	CA	HIS A	18		1.495	-4.331	36.754	1.00 54.39
	ATOM	149	C	HIS A	18		1.960	-4.115	35.313	1.00 55.01
	ATOM	150	Ö	HIS A	18		1.722	-3.036	34.757	1.00 54.91
10	ATOM	151	СВ	HIS A	18	•	2.663	-4.168	37.735	1.00 53.19
10	ATOM	152	CG	HIS A	18		2.198	-3.867	39.130	1.00 52.86
		153		HIS A	18		1.486	-2.733	39.432	1.00 52.60
	ATOM	154		HIS A	18		2.362	-4.553	40.296	1.00 52.72
	ATOM				18		1.216	-2.720	40.730	1.00 53.99
1.5	MOTA	155		HIS A			1.735	-3.817	41.269	1.00 53.27
15	MOTA	156	NE2	HIS A	18		2.580	-5.122	34.721	1.00 56.14
	MOTA	157	N	TYR A	19		3.044	-5.034	33.337	1.00 58.17
-	MOTA	158	CA	TYR A	19				32.380	1.00 58.92
	ATOM	159	C	TYR A	19		1.890	-4.733	31.552	1.00 59.36
	ATOM	160	0	TYR A	19		1.957	-3.827		1.00 59.47
20	ATOM	161	CB	TYR A	19		3.759	-6.320	32.950	
	ATOM	162	CG	TYR A	19		5.097	-6.621	33.580	
	ATOM	163		TYR A	19		5.983	-5.607	33.934	1.00 63.22
	MOTA	164	CD2	TYR A	19		5.513	-7.934	33.787	1.00 62.96
	ATOM	165	CE1	TYR A	19		7.212	-5.891	34.488	1.00 63.58
25	ATOM	166	CE2	TYR A	19		6.745	-8.226	34.345	1.00 63.59
	ATOM	167	CZ	TYR A	. 19		7.597	-7.199	34.703	1.00 63.46
	MOTA	168	OH	TYR A	19		8.828	-7.470	35.274	1.00 62.56
	MOTA	169	N	GLU A	20		0.779	-5.451	32.499	1.00 59.63
	ATOM	170	CA	GLU A	20		-0.426	-5.196	31.734	1.00 59.76
30	ATOM	171	С	GLU A	20		-0.990	-3.804	31.918	1.00 59.59
	ATOM	172	0	GLU A	20		-1.198	-3.103	30.928	1.00 59.90
	ATOM	173	CB	GLU A	20		-1.499	-6.241	32.056	1.00 66.29
	ATOM	174	CG	GLU A	20		-1.176	-7.583	31.409	1.00 73.88
	ATOM	175	CD	GLU A	20		-2.142	-8.678	31.815	1.00 78.60
35	ATOM	176	OE1	GLU A	20		-1.749	-9.862	31.692	1.00 82.52
55	ATOM	177	OE2	GLU A	20		-3.272	-8.365	32.242	1.00 78.94
	ATOM	178	N	ASN A	21		-1.186	-3.340	33.145	1.00 59.04
	ATOM	179	CA	ASN A	21		-1.784	-2.053	33.404	1.00 57.97
	ATOM	180	C	ASN A	21		-1.002	-0.853	32.918	1.00 57.50
40	ATOM	181	Õ	ASN A	21		-1.637	0.118	32.496	1.00 57.19
40		182	CB	ASN A	21		-2.149	-1.876	34.875	1.00 61.71
	ATOM		CG	ASN A	21		-3.089	-2.964	35.362	1.00 63.07
	ATOM	183		ASN A	21		-3.691	-3.685	34.563	1.00 60.88
	ATOM	184			21		-3.161	-3.066	36.685	1.00 62.59
45	ATOM	185		ASN A			0.327	-0.826	33.022	1.00 56.38
43	ATOM	186	N	ASP A	22 22		1.080	0.299	32.480	1.00 54.54
	ATOM	187	CA	ASP A		•	0.710	1.630	33.112	1.00 53.00
	ATOM	188	C	ASP A	22		0.632	2.652	32.424	1.00 52.65
	ATOM	189	0	ASP A	22			0.369	30.967	1.00 61.62
50	MOTA	190	CB	ASP A	22		0.818	0.655	30.214	1.00 65.99
50	ATOM	191	CG	ASP A	22		2.107		30.214	1.00 64.95
	ATOM	192		ASP A	22		2.946	1.402	30.765	1.00 69.58
	ATOM	193		ASP A	22		2.235	0.116	29.099	
	ATOM	194	N	SER A	23		0.616	1.683	34.440	1.00 50.98
	ATOM	195	CA	SER A	23		0.213	2.879	35.148	1.00 49.25
55	MOTA	196	С	SER A	23		1.294	3.960	35.095	1.00 48.02
	ATOM	197	0	SER A	23		2.450	3.746	34.740	1.00 47.57
	ATOM	198	CB	SER A	23		-0.101	2.633	36.640	1.00 46.04
	ATOM	199	OG	SER A	23		0.549	1.424	36.984	1.00 56.16
	ATOM	200	N	THR A	24		0.847	5.152	35.441	1.00 47.42
60	ATOM	201	CA	THR A	24		1.724	6.312	35.512	1.00 48.00
	ATOM	202	С	THR A	24		1.731	6.860	36.920	1.00 47.34
	ATOM	203	0	THR A	24		2.328	7.898	37.173	1.00 47.41
	ATOM	204	СВ	THR A	24		1.369	7.421	34.505	1.00 50.50

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	ATOM	205	OG1	THR A	24	•	0.042	7.871	34.734		51.02
	ATOM	206	CG2	THR A	24		1.558	6.901	33.094		48.60
	ATOM	207	N	ASP A	25		1.124	6.096	37.828		46.98
	ATOM	208	CA	ASP A	25		1.058	6.453	39.234	1.00	46.50
5	ATOM	209	С	ASP A	25		2.193	5.788	40.013		45.63
•	ATOM	210	0	ASP A	25		2.376	4.578	40.042	1.00	44.95
	ATOM	211	СB	ASP A	25	-	0.286	6.024	39.847	1.00	53.36
	ATOM	212	CG	ASP A	25	_	1.442	6.789	39.202	1.00	62.10
	ATOM	213		ASP A	25		1.605	7.997	39.498	1.00	64.72
10	ATOM	214		ASP A	25		2.185	6.192	38.392	1.00	62.72
10	ATOM	215	N	LEU A	26		3.000	6.633	40.614	1.00	45.40
	ATOM	216	CA	LEU A	26		4.167	6.207	41.381		45.37
		217	C	LEU A	26		3.834	5.157	42.418		45.85
	ATOM			LEU A	26		4.563	4.170	42.565		46.38
15	ATOM	218	0		26		4.763	7.483	41.982	1.00	44.87
15	ATOM	219	CB	LEU A			6.056	7.341	42.783	1.00	44.69
	ATOM	220	CG	LEU A	26				41.931	1.00	40.20
	MOTA	221		LEU A	26		7.128	6.688		1.00	46.93
	ATOM	222		LEU A	26		6.529	8.703	43.267		
	MOTA	223	N	ARG A	27		2.741	5.281	43.178	1.00	45.28
20	MOTA	224	CA	ARG A	27		2.266	4.241	44.065	1.00	45.19
	MOTA	225	С	ARG A	<b>27</b> .		2.251	2.841	43.483		44.47
	ATOM	226	0	ARG A	27		2.610	1.886	44.187		44.56
	ATOM	227	CB	ARG A	27		0.852	4.494	44.607		51.04
	MOTA	228	CG	ARG A	27		0.713	5.531	45.690		61.27
25	MOTA	229	CD	ARG A	27	-	0.715	6.081	45.714		66.89
	ATOM	230	NE	ARG A	27	-	0.927	6.984	46.839		75.54
	ATOM	231	CZ	ARG A	27	-	2.083	7.555	47.170	1.00	79.36
	ATOM	232		ARG A	27	_	3.184	7.331	46.456		80.70
	ATOM	233		ARG A	27	_	2.152	8.359	48.228	1.00	79.78
30	ATOM	234	N	ASP A	28		1.771	2.632	42.255	1.00	43.06
-	ATOM	235	CA	ASP A	28		1.785	1.318	41.648	1.00	41.77
	ATOM	236	C	ASP A	28		3.195	0.797	41.385	1.00	41.18
	MOTA	237	ō	ASP A	28		3.408	-0.409	41.402	1.00	40.08
	ATOM	238	СВ	ASP A	28		1.014	1.325	40.303	1.00	44.91
35	ATOM	239	CG	ASP A	28	_	0.450	1.655	40.560	1.00	53.31
55		240		ASP A	28		1.014	2.560	39.920		53.29
	ATOM			ASP A	28		1.022	0.983	41.446		52.80
	ATOM	241			29		4.132	1.700	41.062		40.87
	ATOM	242	И	HIS A			5.510	1.269	40.787		40.10
40	ATOM	243	CA	HIS A	29 29		6.208	0.795	42.073	1.00	39.08
40	ATOM	244	C	HIS A				-0.143	42.045	1.00	38.12
	ATOM	245	0	HIS A	29		6.987		40.166		40.60
	ATOM	246	CB	HIS A	29		6.246	2.473			42.13
	MOTA	247	CG	HIS A	29		5.590	2.806	38.837		42.28
	MOTA	248		HIS A	29		5.069	1.810	38.042	-	
45	MOTA	249		HIS A	29		5.373	3.980	38.192		44.10
	MOTA	250		HIS A	29		4.552	2,348	36.943		43.73
	MOTA	251	NE2	HIS A	29		4.738	3.656	37.014		42.95
	MOTA	252	И	ILE A	30		5.896	1.454	43.152	1.00	39.44
	MOTA	253	CA	ILE A	30		6.339	0.990	44.501		39.95
50	ATOM	254	С	ILE A	30		5.899	-0.426	44.746		40.09
	MOTA	255	0	ILE A	30		6.658	-1.303	45.181		41.16
	MOTA	256	CB	ILE A	30		5.843	1.991	45.550	1.00	40.65
	MOTA	257		ILE A	30		6.563	3.321	45.334	1.00	40.86
	ATOM	258		ILE A	30		6.125	1.537	47.004	1.00	41.39
55	ATOM	259		ILE A	30		6.060	4.498	46.138	1.00	42.24
-	ATOM	260	N	ASP A	31		4.631	-0.764	44.485	1.00	41.09
	ATOM	261	CA	ASP A	31		4.082	-2.089	44.758		40.37
	•	262	C	ASP A	31		4.718	-3.130	43.856		40.59
	ATOM ATOM	263	0	ASP A	31		4.965	-4.277	44.244		40.70
60		264		ASP A	31		2.566	-2.080	44.459		42.53
UU	ATOM		CB		31		1.886	-3.379	44.801		44.66
	ATOM	265	CG	ASP A			1.799	-4.311	43.991		46.03
	ATOM	266		ASP A	31			-3.517	45.987		53.28
	ATOM	267	002	ASP A	31		1.495	-3.31/	33.307	1.00	JJ.20

					20	4 045	0 775	42.589	1.00 39.00
	ATOM	268	N	TYR A	32	4.945	-2.735		1.00 39.00
	ATOM	269	CA	TYR A	32	5.636	-3.647	41.677	1.00 36.55
	ATOM	270	С	TYR A	32	7.017	~4.030	42.231	1.00 36.33
_	MOTA	271	0_	TYR A	32	7.359	-5.204	42.252 40.324	1.00 30.44
5	ATOM	272	CB	TYR A	32	5.765	-2.921	39.369	1.00 33.37
	ATOM	273	CG	TYR A	32	6.750	-3.532		1.00 45.04
	ATOM	274	CD1	TYR A	32	6.374	-4.668	38.646	1.00 43.04
	ATOM	275	CD2	TYR A	32	8.005	-2.989	39.141	1.00 46.06
10	ATOM	276	CE1	TYR A	32	7.245	-5.272	37.758 38.235	1.00 44.10
10	MOTA	277	CE2	TYR A	32	8.871 8.489	-3.576 -4.707	37.545	1.00 45.75
	ATOM	278	CZ	TYR A	32 32	9.322	-5.303	36.633	1.00 44.58
	MOTA	279	OH	TYR A	33	7.850	-3.064	42.552	1.00 36.14
	ATOM	280	N	TRP A	33	9.183	-3.384	43.061	1.00 36.59
15	MOTA	281 282	CA C	TRP A	33	9.144	-4.146	44.391	1.00 36.80
15	ATOM ATOM	283	ŏ	TRP A	33	10.050	-4.951	44.634	1.00 37.42
	ATOM	284	CB	TRP A	33	10.054	-2.131	43.159	1.00 37.44
	ATOM	285	CG	TRP A	33	10.588	-1.813	41.780	1.00 34.77
	ATOM	286		TRP A	33	10.244	-0.745	40.979	1.00 35.47
20	ATOM	287	CD2	TRP A	33	11.522	-2.605	41.047	1.00 32.86
20	ATOM	288	NE1	TRP A	33	10.974	-0.822	39.805	1.00 32.84
	ATOM	289	CE2	TRP A	33	11.735	-1.947	39.799	1.00 35.43
	ATOM	290	CE3	TRP A	33	12.209	-3.792	41.301	1.00 32.38
	ATOM	291	CZ2	TRP A	33	12.595	-2.444	38.832	1.00 35.55
25	ATOM	292	CZ3	TRP A	33	13.061	-4.282	40.337	1.00 37.27
	ATOM	293	CH2	TRP A	33	13.245	-3.626	39.108	1.00 38.28
	ATOM	294	N	LYS A	34	8.150	-3.912	45.246	1.00 37.05
	ATOM	295	CA	LYS A	34	7.990	-4.765	46.437	1.00 37.56
	ATOM	296	С	LYS A	34	7.687	-6.205	46.060	1.00 37.90
30	ATOM	297	0	LYS A	34	8.220	-7.124	46.684	1.00 37.29
	MOTA	298	CB	LYS A	34	6.860	-4.261	47.345	1.00 36.29
	ATOM	299	CG	LYS A	34	7.111	-2.871	47.891	1.00 41.05
	MOTA	300	CD	LYS A	34	6.095	-2.498	48.945	1.00 47.20
	ATOM	301	CE	LYS A	34	5.764	-1.032	48.964	1.00 49.66
35	MOTA	302	NZ	LYS A	34	5.046	-0.625	50.219	1.00 57.26 1.00 37.50
	ATOM	303	N	HIS A	35	6.853	-6.411	45.025	
	MOTA	304	CA	HIS A	35	6.670	-7.779	44.525	1.00 36.43 1.00 35.96
	ATOM	305	C	HIS A	35	7.913	-8.328	43.875 43.986	1.00 34.61
40	ATOM	306	0	HIS A	35	8.237	-9.523 -7.901	43.587	1.00 40.23
40	ATOM	307	CB	HIS A	35 35	5.446 4.200	~7.883	44.428	1.00 44.09
	ATOM	308	CG	HIS A	35	3.567	-6.711	44.788	1.00 48.39
	ATOM	309		HIS A	35	3.539	-8.879	45.058	1.00 48.71
	ATOM ATOM	310 311		HIS A	35	2.538	-6.985	45.574	1.00 48.93
45	ATOM	312		HIS A	35	2.524	-8.283	45.774	1.00 47.98
73	ATOM	313	N	MET A	36	8.665	-7.457	43.180	1.00 35.13
	ATOM	314	CA	MET A	36	9.927	-7.985	42.606	1.00 35.12
	ATOM	315	C	MET A	36	10.836	-8.474	43.753	1.00 35.05
	ATOM	316	ō	MET A	36	11.472	-9.504	43.634	1.00 34.78
50	ATOM	317	CB	MET A		10.584	-6.890	41.772	1.00 36.78
	ATOM	318	CG	MET A	36	9.832	-6.601	40.454	1.00 38.38
	ATOM	319	SD	MET A	36	10.026	-7.870	39.206	1.00 39.79
	ATOM	320	CE	MET A	36	11.681	-7.505	38.605	1.00 43.43
	ATOM	321	N	ARG A	37	10.903	-7.746	44.853	1.00 35.39
55	ATOM	322	CA	ARG A	37	11.729	-8.145	46.004	1.00 35.09
	MOTA	323	С	ARG A	37	11.240	-9.438	46.667	1.00 34.61
	ATOM	324	0	ARG A	37		-10.319	46.996	1.00 33.53
	MOTA	325	CB	ARG A	37	11.555	~7.001	47.018	1.00 34.72
	ATOM	326	CG	ARG A	37	12.370	-7.186	48.305	1.00 34.13
60	MOTA	327	CD	ARG A	37	12.132	-5.981	49.197	1.00 34.07
	ATOM	328	NE	ARG A	37	12.665	-6.189	50.551	1.00 35.48
	MOTA	329	CZ	ARG A	37	12.420	-5.313	51.520	1.00 33.64
	ATOM	330	NH1	ARG A	37	11.676	-4.228	51.375	1.00 38.49

					~-	10 040 5 572	E0 710	1.00 32.86
	MOTA	331		ARG A	37	12.948 -5.572	52.719	1.00 32.80
	ATOM	332	N	LEU A	38	9.920 -9.544	46.841 47.372	1.00 34.82
	ATOM	333	CA	LEU A	38	9.330 -10.768	46.532	1.00 34.02
_	ATOM	334	C	LEU A	38	9.592 -11.985 9.879 -13.050	40.332	1.00 35.88
5	MOTA	335	0	LEU A	38	7.806 -10.549	47.542	1.00 37.00
	ATOM	336	CB	LEU A	38	7.048 -11.843	47.862	1.00 39.41
	ATOM	337	CG	LEU A	38	7.338 -12.303	49.271	1.00 36.17
	ATOM	338		LEU A	38 38	5.544 -11.698	47.633	1.00 42.91
10	ATOM	339		LEU A	39	9.532 -11.873	45.176	1.00 36.01
10	ATOM	340	N	GLU A	39	9.903 -12.982	44.328	1.00 35.41
	ATOM	341	CA	GLU A GLU A	39	11.310 -13.492	44.650	1.00 35.95
	ATOM	342	C	GLU A	39	11.524 -14.706	44.610	1.00 34.71
	ATOM	343	O CB	GLU A	39 .	9.826 -12.621	42.814	1.00 33.83
15	ATOM	344 345	CG	GLU A	39	9.999 -13.858	41.944	1.00 35.55
1.5	MOTA	346	CD	GLU A	39	10.153 -13.499	40.467	1.00 44.56
	ATOM ATOM	347		GLU A	39	11.229 -12.997	40.106	1.00 42.84
	ATOM	348		GLU A	39	9.219 -13.700	39.690	1.00 42.80
	ATOM	349	N	CYS A	40	12.280 -12.600	44.916	1.00 35.37
20	MOTA	350	CA	CYS A	40	13.616 -13.054	45.262	1.00 35.02
20	ATOM	351	C	CYS A	40	13.603 -13.852	46.574	1.00 35.78
	ATOM	352	ŏ	CYS A	40	14.329 -14.842	46.621	1.00 35.14
	MOTA	353	СВ	CYS A	40	14.587 -11.879	45.434	1.00 34.19
	ATOM	354	SG	CYS A	40	14.743 -10.845	43.945	1.00 35.07
25	ATOM	355	N	ALA A	41	12.796 -13.419	47.540	1.00 36.59
	ATOM	356	CA	ALA A	41	12.772 -14.160	48.820	1.00 38.13
	ATOM	357	C	ALA A	41	12.191 -15.553	48.590	1.00 37.52
	ATOM	358	Ō	ALA A	41	12.659 -16.527	49.200	1.00 38.32
	ATOM	359	CB	ALA A	41	11.955 -13.380	49.827	1.00 36.08
30	ATOM	360	N	ILE A	42	11.221 -15.674	47.663	1.00 37.54
	ATOM .	361	CA	ILE A	42	10.629 -16.995	47.397	1.00 36.18
	ATOM	362	С	ILE A	42	11.626 -17.922	46.753.	1.00 36.44
	ATOM	363	0	ILE A	42	11.856 -19.069	47.133	1.00 35.31
	MOTA	364	CB	ILE A	42	9.325 -16.907	46.581	1.00 36.30
35	ATOM	365	CG1	ILE A	42	8.225 -16.165	47.345	1.00 38.51
	MOTA	366	CG2		42	8.865 -18.282	46.108	1.00 38.23
	MOTA	367	CD1	ILE A	42	7.114 -15.700	46.390	1.00 41.57
	ATOM	368	N	TYR A	43	12.321 -17.436	45.707	1.00 35.53
	ATOM	369	CA	TYR A	43	13.341 ~18.254	45.060	1.00 36.20
40	ATOM	370	C	TYR A	43	14.479 -18.536	46.047	1.00 36.21
	ATOM	371	O ·	TYR A	43	15.091 ~19.597	45.993	1.00 36.69 1.00 36.61
	ATOM	372	CB	TYR A	43	13.884 ~17.474	43.838 42.572	1.00 38.55
	ATOM	373	CG	TYR A	43	13.065 -17.637 12.717 -16.512	41.820	1.00 30.55
15	MOTA	374		TYR A	43	12.717 -16.512 12.644 -18.871	42.116	1.00 39.53
45	ATOM	375	CD2	TYR A	43 43	11.998 -16.626	40.646	1.00 41.22
	ATOM	376 377	CE2		43	11.943 -19.006	40.918	1.00 40.40
	ATOM ATOM	378	CZ	TYR A	43	11.604 -17.884	40.210	1.00 40.78
	ATOM	379	OH	TYR A	43	10.847 -17.954	39.081	1.00 41.19
50	MOTA	380	N	TYR A	44	14.794 -17.563	46.906	1.00 35.43
50	ATOM	381	CA	TYR A	44	15.933 -17.815	47.811	1.00 37.45
	MOTA	382	C	TYR A	44	15.547 -19.008	48.716	1.00 37.65
	MOTA	383	ŏ	TYR A	44	16.329 -19.945	48.876	1.00 38.00
		384	СB	TYR A	44	16.205 -16.555	48.635	1.00 38.20
55	ATOM	385	CG	TYR A	44	17.445 -16.670	49.503	1.00 40.09
	ATOM	386		TYR A	44	17.398 -17.286	50.756	1.00 41.30
	ATOM	387		TYR A	44	18.663 -16.206	49.041	1.00 40.77
	ATOM	388		TYR A	44	18.569 -17.412	51.492	1.00 42.23
	MOTA	389		TYR A	44	19.833 -16.312	49.776	1.00 42.80
60	ATOM	390	CZ	TYR A	44	19.746 -16.907	51.023	1.00 43.39
	ATOM	391	OH	TYR A	44	20.863 -17.049	51.798	1.00 45.51
	ATOM	392	N	LYS A	45	14.334 -18.982	49.224	1.00 38.11
	ATOM	393	CA	LYS A	45	13.891 -20.078	50.118	1.00 40.98

	ATOM ATOM	394 395	C O	LYS LYS		45 45	13.832 -21.403 14.315 -22.472	49.387 49.789		41.49 42.02
	MOTA	396	CB	LYS		45	12.537 -19.754	50.750		40.25
5	ATOM	397	CG	LYS		45	11.968 -20.894	51.614		48.19
3	ATOM ATOM	398 399	CD	LYS LYS		45 45	12.824 -21.269 12.671 -20.308	52.813 53.983		51.93 59.68
	ATOM	400	CE NZ	LYS		45	13.979 -20.145	54.698		61.34
	ATOM	401	N	ALA		46	13.307 -21.357	48.139		42.39
	ATOM	402	CA	ALA		46	13.230 -22.586	47.356		41.88
10	MOTA	403	С	ALA	A	46	14.613 -23.167	47.179	1.00	41.96
	ATOM	404	0	ALA		46	14.828 -24.368	47.347		42.06
	ATOM	405	CB	ALA		46	12.561 -22.294	46.004		45.41
	ATOM	406	N	ARG		47	15.605 -22.341	46.839		42.18
15	MOTA MOTA	407 408	CA C	ARG ARG		47 47	16.967 -22.806 17.567 -23.364	46.649 47.949		42.89 44.97
13	ATOM	409	Ö	ARG		47	18.270 -24.377	47.899	1.00	
	ATOM	410	СВ	ARG		47	17.873 -21.700	46.134	1.00	
	ATOM	411	CG	ARG		47	19.278 -22.115	45.751	1.00	
	ATOM	412	CD	ARG		47	19.323 -23.087	44.564		51.87
20	ATOM	413	NE	ARG		47	20.701 -23.450	44.306	1.00	57.92
	MOTA	414	CZ	ARG		47	21.372 -24.042	43.351	1.00	
	ATOM	415	NH1			47	20.819 -24.506	42.243	1.00	
	ATOM	416	NH2			47	22.696 -24.175	43.516	1.00	
25	ATOM ATOM	417 418	N CA	GLU GLU		48 48	17.287 -22.673 17.783 -23.090	49.051 50.364	1.00	45.14 48.18
23	ATOM	419	C	GLU		48	17.266 -24.500	50.681		48.66
	ATOM	420	ō	GLU		48	18.009 -25.362	51.122		49.01
	ATOM	421	CB	GLU		48	17.202 -22.208	51.472		50.97
	ATOM	422	CG	GLU	Α	48	17.987 -21.039	51.955	1.00	59.37
30	ATOM	423	CD	GLU		48	17.911 -20.689	53.432		58.91
	ATOM	424	OE1			48	16.891 -20.144	53.912		64.65
	ATOM	425		GLU		48	18.904 -20.949	54.156		64.43
	ATOM ATOM	426 427	N CA	MET MET		49 49	16.000 -24.730 15.358 -26.017	50.361 50.510		49.75 50.42
35	ATOM	428	C	MET		49	15.738 -27.106	49.543		50.69
	ATOM	429	ŏ	MET		49	15.197 -28.218	49.620		51.52
	ATOM	430	СВ	MET		49	13.840 -25.835	50.513		53.09
	MOTA	431	CG		A	49	13.351 -25.048	51.719	1.00	55.67
40	ATOM	432	SD	MET		49	11.617 -24.613	51.508		64.13
40	ATOM	433	CE	MET		49	10.922 -24.927	53.123		64.34
	ATOM ATOM	434 435	N CA	GLY GLY		50 50	16.616 -26.852 17.159 -27.787	48.592 47.664		50.09 48.51
	ATOM	435	C	GLY		50	16.332 -27.949	46.384		48.34
	ATOM	437	Ö	GLY		50	16.603 -28.920	45.679		47.16
45	ATOM	438	N	PHE		51	15.383 -27.074	46.084		48.32
	MOTA	439	CA	PHE .		51	14.628 -27.205	44.837	1.00	48.99
	ATOM	440	С	PHE .		51	15.442 -26.616	43.675		49.10
	ATOM	441	0	PHE.		51	16.187 -25.659	43.884		48.66
50	ATOM	442	CB	PHE .		51	13.266 -26.527	44.904		54.66
30	ATOM ATOM	443 444	CG	PHE .		51 51	12.370 -27.030 12.237 -28.382	46.005 46.261		62.30 66.64
	ATOM	445		PHE		51	11.648 -26.142	46.784		64.64
	ATOM	446		PHE		51	11.418 -28.837	47.281		67.20
	ATOM	447		PHE		51	10.813 -26.584	47.788		65.80
55	ATOM	448	CZ	PHE A		51	10.708 -27.937	48.047		65.88
	MOTA	449	N	LYS 2	A	52	15.455 -27.331	42.554		49.49
	ATOM	450	CA	LYS I		52	16.204 -26.877	41.377		48.93
	MOTA	451	C	LYS A	A.	52	15.254 -26.192	40.393		47.94
60	ATOM ATOM	452		LYS A		52	15.644 -25.414	39.516		46.65
JU	ATOM	453 454	CB CG	LYS A		52 52	16.905 -28.073 17.964 -28.697	40.739 41.640		54.00 61.04
	ATOM	455		LYS A		52	18.931 -29.536	40.811		64.27
	ATOM	456		LYS A		52	19.717 -30.537	41.711	0.00	

	- mos/	457	117	LYS A	52	20.300 -31.674 41.001 0.00 9	9.00
	ATOM ATOM	457 458	NZ N	HIS A	53	13.968 -26.480 40.589 1.00 4	
	ATOM	459	CA	HIS A	53	12.924 -25.816 39.834 1.00 4	
	ATOM	460	C	HIS A	53		4.64
5	ATOM	461	ŏ	HIS A	53	11.566 -26.465 41.649 1.00 4	4.95
,	ATOM	462	CB	HIS A	53		4.28
	ATOM	463	CG	HIS A	53	11.635 -27.700 38.758 1.00 4	6.45
	ATOM	464		HIS A	53	10.277 -27.525 38.978 1.00 4	15.21
	ATOM	465		HIS A	53	11.868 -29.026 38.829 1.00 4	15.27
10	ATOM	466		HIS A	53	9.709 -28.698 39.163 1.00 4	
	ATOM	467		HIS A	53	10.655 -29.629 39.076 1.00 4	
	ATOM	468	N	ILE A	54	10.895 -24.693 40.468 1.00 4	
	ATOM	469	CA	ILE A	54	9.598 -24.513 41.100 1.00 4	
	ATOM	470	С	ILE A	54	8.589 -24.354 39.976 1.00 4	
15	MOTA	471	0	ILE A	54	8.704 -23.454 39.123 1.00 4	
	ATOM	472	CB	ILE A	54	9.627 -23.258 41.996 1.00 5	
	MOTA	473	CG1	ILE A	54	8.222 -22.740 42.337 1.00 5	
	ATOM	474	CG2	ILE A	54	10.420 -22.138 41.313 1.00 5	
	ATOM	475	CD1	ILE A	54	8.293 -22.022 43.690 1.00 5	
20	MOTA	476	N	ASN A	55	7.679 -25.327 39.853 1.00 4	
	ATOM	477	CA	ASN A	55	6.719 -25.332 38.748 1.00 4	
	ATOM	478	С	ASN A	55	7.391 -25.383 37.389 1.00 4	
	MOTA	479	0	ASN A	55	6.958 -24.848 36.368 1.00 4	
	MOTA	480	CB	ASN A	55	5.639 -24.266 38.828 1.00 4	
25	ATOM	481	CG	ASN A	55	4.611 -24.596 39.915 1.00 4	
	MOTA	482		ASN A	55	4.481 -25.766 40.285 1.00 4	
	ATOM	483		ASN A	55	3.908 -23.589 40.393 1.00 4 8.508 -26.115 37.343 1.00 4	
	ATOM	484	N	HIS A	56		
	ATOM	485	CA	HIS A	56	9.304 -26.336 36.155 1.00 4 9.996 -25.085 35.651 1.00 4	
30	ATOM	486	C	HIS A	56		
	ATOM	487	0	HIS A	56	10.579 -25.115 34.573 1.00 4 8.509 -27.081 35.059 1.00 3	
	ATOM	488	CB	HIS A	56		
	ATOM	489	CG	HIS A	56	8.140 -28.417 35.639 1.00 3 8.997 -29.476 35.683 1.00 4	
25	ATOM	490		HIS A	56 56	7.000 -28.811 36.253 1.00 4	
35	ATOM	491		HIS A	56		16.16
	ATOM	492		HIS A	56	7.192 -30.108 36.648 1.00 4	
	ATOM	493		HIS A GLN A	57	10.115 -24.074 36.492 1.00 4	
	ATOM	494 495	N CA		57	10.867 -22.845 36.209 1.00	
40	ATOM	495	CA	GLN A	57	12.178 -22.922 37.011 1.00	
40	ATOM ATOM	497	Ö	GLN A	57	12.135 -23.367 38.159 1.00 4	
	ATOM	498	СВ	GLN A	57	10.017 -21.730 36.771 1.00	14.03
	ATOM	499	CG	GLN A	57	10.191 -20.286 36.476 1.00 5	54.62
	ATOM	500	CD	GLN A	57	8.889 -19.530 36.723 1.00 5	
45	ATOM	501		GLN A	57	8.881 -18.332 36.990 1.00 5	54.35
	ATOM	502		GLN A	57	7.774 -20.248 36.641 1.00 5	
	ATOM	503	N	VAL A	58	13.308 -22.526 36.435 1.00 4	
	ATOM	504		VAL A	58	14.592 -22.695 37.122 1.00 4	
	ATOM	505	С	VAL A	58	14.654 -21.870 38.395 1.00	39.90
50	ATOM	506	0	VAL A	58	14.141 -20.751 38.425 1.00	
	ATOM	507	CB	VAL A	58	15.766 -22.343 36.168 1.00	
	ATOM	508	CG1	VAL A	58	15.770 -20.860 35.859 1.00	
	ATOM	509	CG2	VAL A	58	17.085 -22.872 36.676 1.00	
	ATOM	510	N	VAL A	59	15.193 -22.451 39.461 1.00	
55	ATOM	511	CA	VAL A	59	15.414 -21.673 40.713 1.00	
	ATOM	512	-C	VAL A	59	16.878 -21.205 40.635 1.00	
	ATOM	513	0	VAL A	59	17.761 -22.042 40.559 1.00	
	ATOM	514	CB	VAL A	59	15.250 -22.591 41.945 1.00	
	ATOM	515		VAL A	59	15.437 -21.777 43.243 1.00	
60	MOTA	516		VAL A	59	13.830 -23.161 42.037 1.00 3	
	ATOM	517	N	PRO A	60	17.121 -19.911 40.686 1.00	
	MOTA	518	CA	PRO A	60	18.466 -19.380 40.576 1.00	
	ATOM	519	С	PRO A	60	19.312 -19.737 41.806 1.00	10.5/

5	MOTA MOTA MOTA	520 521 522 523	O CB CG CD	PRO A PRO A PRO A	60 60 60 61	18.766 -20.039 18.272 -17.874 16.837 -17.576 16.069 -18.866 20.627 -19.662	42.863 40.580 40.657 40.722 41.632	1.00 39.38 1.00 40.12 1.00 40.26 1.00 39.33 1.00 39.59
5	ATOM ATOM ATOM ATOM ATOM	524 525 526 527 528	N CA C O CB	THR A THR A THR A THR A	61 61 61	21.502 -19.731 21.145 -18.651 20.503 -17.621 23.000 -19.670	42.799 43.803 43.545 42.486	1.00 39.51 1.00 38.37 1.00 37.40 1.00 42.80
10	ATOM ATOM ATOM ATOM ATOM	529 530 531 532 533	OG1 CG2 N CA C		61 61 62 62 62	23.259 -18.498 23.435 -20.914 21.544 -18.931 21.280 -17.966 21.841 -16.582	41.695 41.722 45.058 46.132 45.847	1.00 42.80 1.00 42.91 1.00 37.77 1.00 36.91 1.00 36.03
15	ATOM ATOM ATOM ATOM	534 535 536 537	O CB CG CD1	LEU A LEU A LEU A	62 62 62 62	21.248 -15.566 21.904 -18.492 21.278 -19.807 22.090 -20.339	46.209 47.447 47.950 49.139	1.00 36.36 1.00 35.93 1.00 36.47 1.00 44.24
20	ATOM ATOM ATOM ATOM ATOM	538 539 540 541 542	CD2 N CA C	LEU A ALA A ALA A ALA A	62 63 63 63	19.850 -19.557 23.053 -16.511 23.729 -15.236 22.946 -14.476 22.808 -13.263	48.450 45.336 45.091 44.014 44.128	1.00 33.58 1.00 36.59 1.00 36.90 1.00 36.28 1.00 35.61
25	ATOM ATOM ATOM ATOM	543 544 545 546	CB N CA C	ALA A VAL A VAL A	63 64 64 64	25.153 -15.402 22.430 -15.198 21.587 -14.456 20.370 -13.842	44.601 43.013 42.054 42.720	1.00 39.35 1.00 35.11 1.00 35.89 1.00 35.12
30	ATOM ATOM ATOM	547 548 549 550	CG2	VAL A VAL A VAL A	64 64 64 65	20.062 -12.664 21.180 -15.331 20.071 -14.660 22.390 -15.680 19.616 -14.592	42.487 40.855 40.053 40.014 43.540	1.00 34.59 1.00 37.44 1.00 40.96 1.00 36.95 1.00 35.09
35	ATOM ATOM ATOM ATOM ATOM	551 552 553 554 555	N CA C O CB	SER A SER A SER A SER A	65 65 65	18.477 -14.039 18.854 -12.961 18.110 -11.986 17.583 -15.074	44.238 45.225 45.326 44.940	1.00 34.42 1.00 34.48 1.00 34.19 1.00 34.49
40	ATOM ATOM ATOM ATOM	556 557 558 559	OG N CA C	SER A LYS A LYS A LYS A	65 66 66	17.165 -16.015 19.977 -13.079 20.365 -11.976 20.611 -10.670	43.951 45.922 46.828 46.044	1.00 35.80 1.00 34.79 1.00 35.44 1.00 35.50
4.5	ATOM ATOM ATOM ATOM	560 561 562 563	O CB CG CD	LYS A LYS A LYS A	66 66 66	20.219 -9.590 21.709 -12.362 21.492 -13.207 22.772 -13.202	46.478 47.478 48.738 49.570	1.00 35.28 1.00 36.28 1.00 43.87 1.00 52.29
45	ATOM ATOM ATOM	564 565 566 567	CE NZ N CA	LYS A LYS A ASN A ASN A	66 66 67 67	23.722 -14.338 24.326 -14.857 21.345 -10.788 21.679 -9.615	49.256 50.541 44.956 44.136 43.544	1.00 56.53 1.00 60.41 1.00 35.53 1.00 35.43 1.00 34.46
50	ATOM ATOM ATOM ATOM	568 569 570 571	C O CB CG	ASN A ASN A ASN A ASN A	67 67 67 67 67	20.452 -8.955 20.370 -7.741 22.657 -10.003 22.999 -8.797 22.646 -8.711		
55	MOTA ATOM ATOM MOTA ATOM	572 573 574 575 576		ASN A LYS A LYS A LYS A	67 68 68 68	23.611 -7.794 19.505 -9.746 18.245 -9.160 17.421 -8.525	42.784 43.007 42.532 43.620	1.00 51.11 1.00 34.69 1.00 33.90 1.00 33.51
<i>6</i> 0	ATOM ATOM ATOM ATOM	577 578 579 580	O CB CCD	LYS A LYS A LYS A LYS A	68 68 68 68	16.726 -7.530 17.439 -10.259 18.132 -10.664 17.352 -11.710	43.412 41.811 40.503 39.729	1.00 32.16 1.00 35.44 1.00 38.74 1.00 44.24
	ATOM ATOM	581 582	CE	LYS A LYS A	68 68	15.976 -11.178 15.414 -12.058	39.345 38.260	1.00 46.97 1.00 51.68

						17 305	-9.098	44.856	1.00 33.48
	MOTA	583	N	ALA A	69	17.395	-8.440	45.924	1.00 33.17
	MOTA	584	CA	ALA A	69	16.654 17.299	-7.103	46.282	1.00 31.99
	MOTA	585	С	ALA A	69	16.620	-6.105	46.498	1.00 31.80
_	MOTA	586	0	ALA A	69	16.642	-9.315	47.213	1.00 31.02
5	ATOM	587	CB	ALA A	69	18.627	-7.019	46.312	1.00 32.54
	ATOM	588	N	LEU A	70	19.278	-5.715	46.511	1.00 32.22
	ATOM	589	CA	LEU A	70	18.830	-4.668	45.471	1.00 32.04
	ATOM	590	C	LEU A	70		-3.504	45.815	1.00 32.18
10	MOTA	591	0	LEU A	70	18.604 20.800	-5.876	46.425	1.00 35.34
10	ATOM	592	CB	LEU A	70 70	21.431	-6.582	47.652	1.00 41.35
	MOTA	593 594	CG	LEU A	70	22.952	-6.614	47.488	1.00 47.21
	ATOM	595		LEU A	70	21.124	-5.797	48.927	1.00 42.20
	MOTA	596	N N	GLN A	71	18.732	-5.068	44.222	1.00 32.28
15	MOTA MOTA	597	CA	GLN A	71	18.336	-4.184	43.118	1.00 32.66
13	ATOM	598	C	GLN A	71	16.888	-3.744	43.319	1.00 33.12
	ATOM	599	Õ	GLN A	71	16.599	-2.548	43.262	1.00 34.13
	ATOM	600	СВ	GLN A	71	18.506	-4.847	41.767	1.00 32.48
	ATOM	601	CG	GLN A	71	19.933	-5.100	41.321	1.00 36.30
20	ATOM	602	CD	GLN A	71	20.143	-5.912	40.083	1.00 36.39
20	ATOM	603		GLN A	71		-5.642	39.339	1.00 42.50
	ATOM	604		GLN A	71	19.349	-6.917	39.755	1.00 33.68
	ATOM	605	N	ALA A	72	16.008	-4.697	43.668	1.00 32.97
	ATOM	606	CA	ALA A	72	14.624	-4.358	43.963	1.00 32.73
25	ATOM	607	C	ALA A	72	14.529	-3.414	45.129	1.00 33.36
	ATOM	608	Õ	ALA A	72	13.741	-2.468	45.153	1.00 33.02
	ATOM	609	CB	ALA A	72	13.751	-5.597	44.164	1.00 31.63
	ATOM	610	N	ILE A	73	15.314	-3.700	46.205	1.00 32.96
	ATOM	611	CA	ILE A	73	15.341	-2.754	47.321	1.00 32.98
30	ATOM	612	С	ILE A	73	15.756	-1.358	46.932	1.00 33.05
-	ATOM	613	0	ILE A	73	15.173	-0.371	47.407	1.00 31.52
	ATOM	614	CB	ILE A	73	16.262	-3.309	48.450	1.00 32.37
	ATOM	615	CG1	ILE A	73	15.549	-4.497	49.099	1.00 34.48
	ATOM	616	CG2	ILE A	73	16.564	-2.217	49.479	1.00 36.05
35	ATOM	617	CD1	ILE A	73	16.442	-5.452	49.895	1.00 36.56
	ATOM	618	N	GLU A	74	16.821	-1.221	46.107	1.00 33.20
	ATOM	619	CA	GLU A	74	17.249	0.135	45.770	1.00 34.02
	MOTA	620	С	GLU A	74	16.127	0.888	45.042	1.00 33.88
	MOTA	621	0	GLU A	74	15.924	2.077	45.333	1.00 33.58
40	MOTA	622	CB	GLU A	74	18.483	0.128	44.849	1.00 42.88
	MOTA	623	CG	GLU A	74	19.730	-0.391	45.551	1.00 50.60
	MOTA	624	CD	GLU A	74	20.121	0.534	46.697	1.00 55.46
	MOTA	625		GLU A	74	19.809	0.219	47.869 46.386	1.00 51.96
4.5	ATOM	626		GLU A	74	20.627	1.630	44.142	1.00 34.25
45	MOTA	627	N	LEU A	75	15.444	0.203 0.814	43.393	1.00 34.25
	ATOM	628	CA	LEU A	75	14.353 13.181	1.091	44.339	1.00 35.22
	ATOM	629	C	LEU A	75 75	12.683	2.215	44.292	1.00 35.32
	MOTA	630	0.	LEU A	75 75	13.895	-0.038	42.211	1.00 33.68
50	ATOM	631	CB	LEU A		14.632	0.030	40.849	
50	ATOM	632	CG	LEU A	75 75	14.246	1.524	40.263	1.00 38.74
	ATOM	633		LEU A	75	16.134	0.148	41.044	1.00 41.33
		. 634 635		GLN A	76	12.769	0.098	45.129	1.00 35.20
	ATOM		N		76	11.711	0.401	46.107	1.00 34.63
55	ATOM	636 637	CA C	GLN A GLN A	76	12.023	1.622	46.951	1.00 34.20
رر	ATOM	638	0	GLN A	76	11.197	2.539	47.032	1.00 33.10
	ATOM ATOM	639	CB	GLN A	76	11.439	-0.800	47.043	1.00 37.30
	ATOM	640	CG	GLN A	76	10.346	-0.570	48.086	1.00 36.76
	ATOM	641	CD	GLN A	76	10.511	-1.541	49.275	1.00 38.06
60	ATOM	642		GLN A	76	11.019	-2.647	49.179	1.00 36.12
-	MOTA	643		GLN A	76	10.136	-1.178	50.481	1.00 39.68
	MOTA	644	N	LEU A	77	13.195	1.702	47.596	1.00 34.42
	ATOM	645	CA	LEU A	77	13.533	2.857	48.402	1.00 34.29

							_			1.00 34.87
	ATOM	646	С	LEU A			13.506	4.183	47.638	
	ATOM	647	0	LEU A			13.070	5.221	48.149	
	MOTA	648	CB	LEU A			14.906	2.756	49.079	1.00 31.33
	ATOM	649	CG	LEU A			14.976	1.566	50.093	1.00 33.32
5	MOTA	650	CD1	LEU A			16.417	1.466	50.566	1.00 35.24
_	ATOM	651	CD2	LEU A	. 77	'	14.094	1.902	51.303	1.00 32.76
	ATOM	652	N	THR A	. 78	]	14.094	4.162	46.440	1.00 35.43
	ATOM	653	CA	THR A		}	14.147	5.391	45.644	1.00 35.16
	ATOM	654	C	THR A		}	12.754	5.938	45.407	1.00 34.62
10	ATOM	655	ō	THR A		}	12.561	7.128	45.655	1.00 35.48
10	ATOM	656	СВ	THR A			14.869	5.117	44.306	1.00 33.75
	ATOM	657		THR A			16.212	4.853	44.644	1.00 36.55
	ATOM	658					14.710	6.309	43.359	1.00 35.96
		659	N N	LEU A			11.867	5.059	44.971	1.00 35.23
1.5	ATOM		CA	LEU A			10.492	5.458	44.646	1.00 35.59
15	ATOM	660		LEU A			9.738	5.941	45.879	1.00 36.24
	MOTA	661	C				8.923	6.848	45.814	1.00 34.44
	MOTA	662	0	LEU A			9.744	4.326	43.961	1.00 37.56
	MOTA	663	CB	LEU A			10.302	3.825	42.611	1.00 40.81
	ATOM	664	CG	LEU A			9.415	2.708	42.066	1.00 36.86
20	ATOM	665		LEU A				4.981	41.632	1.00 44.33
	MOTA	666	CD2	LEU A			10.404		47.023	
	MOTA	667	N	GLU A			10.058	5.284		1.00 35.25
	MOTA	668	CA	GLU A		)	9.487	5.773	48.285	1.00 35.23
	ATOM	669	С	GLU A	7 8(	)	10.002	7.132	48.672	1.00 35.81
25	ATOM	670	0	GLU P	7 8	)	9.241	7.941	49.182	
	ATOM	671	CB	GLU F	8 (	)	9.805	4.764	49.414	1.00 33.47
	ATOM	672	CG	GLU A	8	)	8.923	3.555	49.368	1.00 36.03
	ATOM	673	CD	GLU F	8 4	0	9.390	2.431	50.293	1.00 39.80
	ATOM	674	OE1	GLU F	8 4	0	10.528	2.453	50.789	1.00 38.34
30	ATOM	675		GLU A		0	8.587	1.482	50.397	1.00 40.54
50	ATOM	676	N	THR A	_	1	11.266	7.474	48.443	1.00 35.85
	ATOM	677	CA	THR A		1	11.759	8.819	48.714	1.00 37.65
	ATOM	678	C	THR A			11.074	9.798	47.742	1.00 39.27
	MOTA	679	ŏ	THR A			10.711	10.894	48.159	1.00 38.93
35	ATOM	680	СВ	THR A			13.277	8.895	48.523	1.00 38.88
33		681	OG1				13.854	8.188	49.626	1.00 41.45
	ATOM	682	CG2		-		13.827	10.315	48.511	1.00 37.76
	ATOM	683	N N	ILE A			10.887	9.360	46.500	1.00 39.69
	ATOM			ILE A			10.176	10.260	45.568	1.00 41.32
40	ATOM	684	CA C	ILE A			8.727	10.458	45.979	1.00 42.17
40	MOTA	685		ILE A			8.195	11.566	45.910	1.00 42.11
	ATOM	686	0	ILE A			10.199	9.735	44.134	1.00 38.27
	ATOM	687	CB	ILE A			11.619	9.500	43.651	1.00 39.22
	ATOM	688	CG1				9.462	10.697	43.194	1.00 37.66
	MOTA	689	CG2		-		12.489	10.717	43.731	1.00 43.59
45	ATOM	690	CD1		_		8.097	9.376	46.426	1.00 43.95
	MOTA	691	Ŋ	TYR	-		6.726	9.469	46.924	1.00 45.07
	MOTA	692	CA	TYR		3		10.496	48.038	1.00 45.84
	MOTA	693	С	TYR A		3	6.597	11.234	48.097	
	MOTA	694	0	TYR A		3	5.613	11.234	47.364	1.00 47.55
50	MOTA	695	CB	TYR I		3	6.229	8.097		1.00 52.04
	MOTA	696	CG	TYR A		3	4.745	8.146	47.683	1.00 53.71
	MOTA	697		TYR 3		3	3.826	8.070	46.643	1.00 53.71
	ATOM	698	CD2	TYR :	A 8	3	4.292	8.292	48.987	1.00 55.75
	ATOM	699		TYR .		3	2.469	8.119	46.899	1.00 55.59
55	ATOM	700		TYR .		3	2.932	8.343	49.245	1.00 55.42
	MOTA	701	CZ	TYR .		3	2.036	8.252	48.199	1.00 56.63
	ATOM	702	ОН	TYR		3	0.691	8.313	48.454	1.00 58.49
	ATOM	703	N	ASN .		4	7.594	10.600	48.932	1.00 45.54
	ATOM	704	CA	ASN		4	7.519	11.621	49.959	1.00 46.22
60	ATOM	705	C	ASN		4	7.965	12.989	49.446	1.00 45.99
00	ATOM	705	ŏ	ASN		4	7.812	13.930	50.226	1.00 46.80
		707	СВ	ASN		4	8.199	11.252	51.257	1.00 45.97
	ATOM	708	CG	ASN		4	7.995	9.970	52.011	1.00 42.27
	ATOM	,00	-0	22014		-				

	MOTA	709	OD1	ASN A	8 4		6.960	9.292	51.955	1.00	48.67
	ATOM	710		ASN A			9.032	9.528	52.753		40.72
	ATOM	711	N	SER A	8.5		8.351	13.244	48.201		44.95
	ATOM	712	CA	SER A			8.830	14.568	47.823		43.57
5	MOTA	713	C	SER A			7.800	15.337	46.988		43.08 41.05
	MOTA	714	0	SER A			6.747 10.108	14.845 14.456	46.627 46.969		43.41
	ATOM	715 716	CB OG	SER A			9.782	14.024	45.656		38.06
	ATOM ATOM	717	N	GLN A			8.199	16.543	46.586		43.58
10	ATOM	718	CA	GLN A			7.377	17.394	45.725	1.00	44.93
	ATOM	719	C	GLN A			7.185	16.853	44.331		45.37
	ATOM	720	0	GLN A	86		6.311	17.326	43.581		45.62
	MOTA	721	CB	GLN A			8.100	18.761	45.588		52.00
	MOTA	722	CG	GLN A			9.448	18.557	44.909 44.692		58.47 64.18
15	ATOM	723	CD	GLN A			10.263 11.330	19.798 19.934	45.303		70.12
	ATOM ATOM	724 725	OE1	GLN A			9.786	20.668	43.810		65.51
	ATOM	726	N .	TYR A			8.002	15.875	43.915		45.13
	ATOM	727	CA	TYR A			7.886	15.277	42.601	1.00	44.09
20	ATOM	728	C	TYR A			6.898	14.125	42.585		44.50
	ATOM .	729	0	TYR A			6.682	13.536	41.536		43.83
	MOTA	730	CB	TYR A			9.231	14.748	42.072		42.80
	MOTA	731	CG	TYR A			10.318	15.782	42.226 43.084		41.15
25	MOTA	732	CD1				11.376 10.272	15.539 17.010	43.064		39.57
25	ATOM	733 734	CD2	TYR A			12.344	16.496	43.319		41.24
	ATOM ATOM	735	CE2	TYR A			11.243	17.967	41.779		39.89
	MOTA	736	CZ	TYR A			12.295	17.693	42.614	1.00	40.85
	ATOM	737	ОН	TYR A			13.302	18.594	42.842		41.44
30	ATOM	738	N	SER A	. 88	t	6.318	13.805	43.743		45.60
	MOTA	739	CA	SER A			5.478	12.626	43.844	-	46.49
	MOTA	740	С	SER A			4.368	12.534	42.814		47.16 46.71
	MOTA	741	0	SER A			4.080 4.816	11.467 12.633	42.270 45.245	-	48.13
35	ATOM	742 743	CB OG	SER A			4.092	11.417	45.346		51.98
33	ATOM ATOM	744	N	ASN A			3.667	13.642	42.565		48.16
	MOTA	745	CA	ASN A			2.563	13.593	41.604	1.00	49.94
	ATOM	746	C	ASN A			2.888	13.746	40.138		50.05
	ATOM	747	0	ASN A	. 89	)	1.922	13.898	39.362		49.81
40	ATOM	748	CB	ASN A			1.500	14.616	42.040		57.17
	MOTA	749	CG	ASN A			1.003	14.286	43.439		62.53 67.35
	ATOM	750	OD1	ASN A			0.752 0.884	15.195 12.995	44.234 43.737		64.39
	ATOM ATOM	751 752	NDZ N	ASN A		, )	4.149	13.696	39.690		49.64
45	ATOM	753	CA	GLU A			4.384	13.669	38.238	-	49.09
15	ATOM	754	C	GLU A			3.952	12.346	37.610		48.60
	ATOM	755	0	GLU A		}	3.715	11.386	38.327		47.76
	ATOM	756	CB	GLU A	. 90	)	5.891	13.782	37.891		47.16
	MOTA	757	CG	GLU A			6.546	14.906	38.632		42.61
50	MOTA	758	CD				7.981	15.219	38.359 37.961		44.84 46.86
	ATOM	759		GLU A			8.767 8.343	14.356 16.400	38.592		45.57
	ATOM ATOM	760 761	N N	GLU A			3.961	12.276	36.269		47.66
	ATOM	762	CA	LYS A			3.798	10.985	35.595		47.27
55	ATOM	763	c	LYS A			5.099	10.180	35.749	1.00	46.58
	ATOM	764	ŏ	LYS A			6.181	10.745	35.610		45.68
	ATOM	765	CB	LYS A			3.615	11.124	34.076		48.52
	ATOM	766	CG	LYS A			2.234	10.887	33.509		55.86
60	MOTA	767	CD	LYS A			2.206 2.934	11.249 10.173	32.017 31.214		56.83 56.89
60	ATOM	768	CE	LYS A			2.934 3.771	10.173	30.132		54.96
	ATOM ATOM	769 770	NZ N	TRP A			4.971	8.888	36.019		46.05
	ATOM	771	CA	TRP A			6.082	7.961	36.101		45.89
	011										

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	ATOM	772	С	TRP A	92	5.702	6.699	35.343	1.00 46.10
	ATOM	773	0	TRP A	92	4.701	6.079	35.668	1.00 46.31
	ATOM	774	CB	TRP A	92	6.454	7.556	37.550	1.00 42.18
	ATOM	775	CG	TRP A	92	6.876	8.733	38.372	1.00 37.28
5	ATOM	776	CD1	TRP A	92	6.025	9.494	39.126	1.00 35.98
	ATOM	777	CD2	TRP A	92	8.170	9.326	38.526	1.00 35.45
	ATOM	778	NE1			6.703	10.512	39.704	1.00 37.68
	ATOM	779	CE2	TRP A	92	8.031	10.426	39.359	1.00 34.31
	ATOM	780	CE3	TRP A		9.436	9.034	38.009	1.00 35.29
10	MOTA	781	CZ2			9.085	11.251	39.784	1.00 37.71
- •	ATOM	782	CZ3	TRP A		10.489	9.844	38.391	1.00 34.81
	ATOM	783		TRP A		10.320	10.920	39.272	1.00 37.49
	ATOM	784	N	THR A		6.468	6.302	34.333	1.00 46.29
	ATOM	785	CA	THR A		6.122	5.105	33.583	1.00 47.56
15	ATOM	786	C	THR A		6.901	3.910	34.110	1.00 48.41
10	ATOM	787	ō	THR A		7.846	4.126	34.862	1.00 48.37
	MOTA	788	СB	THR A		6.370	5.242	32.077	1.00 46.70
	MOTA	789		THR A		7.740	4.928	31.792	1.00 44.41
	ATOM	790	_	THR A		6.037	6.666	31.629	1.00 44.27
20	ATOM	791	N	LEU A	_	6.507	2.716	33.705	1.00 48.40
20		792	CA	LEU A		7.213	1.494	34.076	1.00 48.72
	ATOM	793	C	LEU A		8.647	1.507	33.559	1.00 48.08
	ATOM	794	õ	LEU A		9.587	1.053	34.225	1.00 47.67
	MOTA	795	СВ	LEU A		6.462	0.294	33.479	1.00 56.38
25	MOTA		CG	LEU A		6.488	-1.036	34.232	1.00 61.49
23	MOTA	796				6.452	-0.872	35.741	1.00 61.80
	MOTA	797		LEU A		5.310	-1.900	33.781	1.00 62.20
	ATOM	798		LEU A		8.818	2.022	32.332	1.00 45.95
	ATOM	799	N	GLN A		10.175	2.149	31.805	1.00 43.86
20	ATOM	800		GLN A			3.201	32.555	1.00 41.09
30	MOTA	801	C	GLN A		10.956	2.966	32.803	1.00 42.10
	MOTA	802	0	GLN A		12.138		30.291	1.00 47.51
	MOTA	803	CB	GLN A		10.226	2.431	29.532	1.00 55.82
	MOTA	804	CG	GLN A		10.933	1.311	29.532	1.00 58.57
	MOTA	805	CD	GLN A		12.441	1.369		1.00 63.35
35	ATOM	806		GLN A		13.085	2.191	28.850	1.00 63.33
	ATOM	807		GLN A		13.089	0.470	30.270	1.00 39.88
	MOTA	808	N	ASP A		10.358	4.296	32.984	1.00 39.85
	MOTA	809	CA	ASP A	_	11.048	5.272	33.814	1.00 39.83
	MOTA	810	C	ASP A		11.651	4.615	35.076	1.00 40.22
40	MOTA	811	0	ASP A		12.731	5.042	35,499	
	MOTA	812	CB	ASP A		10.115	6.397	34.246	
	ATOM	813	CG	ASP A		9.603	7.287	33.110	1.00 43.09
	ATOM	814		ASP A		10.290	7.391	32.079	
	ATOM	815		ASP A		8.516	7.883	33.274	
45	MOTA	816	N	VAL A		10.956	3.643	35.651	
	MOTA	817	CA	VAL A		11.371	3.024	. 36.913	1.00 38.84
	MOTA	818	С	VAL A		11.979	1.641	36.744	1.00 38.61
	MOTA	819	0	VAL A		12.059	0.843	37.707	1.00 38.87
	ATOM	820	CB	VAL A		10.186	2.955	37.903	1.00 40.58 1.00 37.24
50	ATOM	821		VAL A		9.561	4.325	38.123	
	ATOM	822	CG2	VAL A		9.122	1.929	37.569	1.00 39.40
	ATOM	823	N	SER A		12.431	1.314	35.539	1.00 37.55
	ATOM	824	CA	SER A		12.956	0.014	35.183	1.00 36.27
1	ATOM	825	С	SER A		14.408	-0.046	35.652	1.00 36.46
55	ATOM	826	0	SER A		15.028	1.015	35.744	1.00 36.95
	ATOM	827	CB	SER A		12.915	-0.142	33.637	1.00 41.66
	ATOM	828	OG	SER A		13.869	0.716	32.996	1.00 41.12
	ATOM	829	N	LEU A	. 99	14.941	-1.247	35.831	1.00 36.38
	ATOM	830	CA	LEU A		16.345	-1.378	36.203	1.00 36.81
60	ATOM	831	С	LEU A	. 99	17.251	-0.939	35.042	1.00 36.83
	ATOM	832	0	LEU A		18.297	-0.341	35.258	1.00 36.14
	MOTA	833	CB	LEU A	. 99	16.636	-2.829	36.589	1.00 42.12
	ATOM	834	CG	LEU A	. 99	18.056	-3.111	37.080	1.00 45.46

								•	•	
	ATOM	835	CD1	LEU A	99	1	18.330	-2.481	38.442	1.00 43.84
	ATOM	836		LEU A			18.341	-4.606	37.072	1.00 45.55
	ATOM	837	N	GLU A			16.797	-1.199	33.818	1.00 36.41
	ATOM	838	CA	GLU A			17.488	-0.765	32.611	1.00 37.21
5	ATOM	839	C	GLU A			17.790	0.722	32.622	1.00 36.02
,		840	Ö	GLU A			18.972	1.068	32.470	1.00 37.81
	ATOM			GLU A			16.690	-1.126	31.340	1.00 40.11
	ATOM	841	CB				17.528	-0.836	30.096	1.00 49.41
	ATOM	842	CG	GLU A					28.790	1.00 56.89
10	MOTA	843	CD	GLU A			16.866	-1.247	28.761	1.00 57.99
10	ATOM	844		GLU A			15.724	-1.755		1.00 57.35
	ATOM	845		GLU A			17.534	-1.025	27.753	
	ATOM	846	N	VAL A		_	16.808	1.580	32.792	1.00 35.41
	ATOM	847	CA	VAL A			17.044	3.010	32.973	1.00 34.54
	ATOM	848	С	VAL A			17.897	3.329	34.185	1.00 34.62
15	MOTA	849	0	VAL A			18.899	4.082	34.151	1.00 33.43
	ATOM	850	CB	VAL A			15.666	3.714	33.047	1.00 36.71
	ATOM	851		VAL A			15.797	5.167	33.459	1.00 34.93
	ATOM	852	CG2	VAL A	101		15.013	3.600	31.651	1.00 37.61
	MOTA	853	N	TYR A	102		17.555	2.678	35.314	1.00 33.28
20	MOTA	854	CA	TYR A	102		18.202	3.008	36.583	1.00 33.58
	ATOM	855	C	TYR A	102	]	19.711	2.817	36.484	1.00 33.46
	ATOM	856	0	TYR A	102	2	20.424	3.624	37.071	1.00 33.95
	ATOM	857	CB	TYR A	102	1	17.653	2.091	37.691	1.00 34.23
	ATOM	858	CG	TYR A	102		17.996	2.494	39.103	1.00 36.50
25	MOTA	859	CD1	TYR A	102	1	17.325	3.559	39.676	1.00 36.36
	ATOM	860	CD2	TYR A			18.961	1.811	39.855	1.00 36.63
	ATOM	861	CE1	TYR A		I	17.582	3.949	40.979	1.00 37.25
	MOTA	862	CE2	TYR A			9.237	2.202	41.162	1.00 37.64
	ATOM	863	CZ	TYR A			8.537	3.253	41.707	1.00 36.95
30	ATOM	864	OH	TYR A	100		8.795	3.693	42.966	1.00 36.81
50	ATOM	865	N	LEU A			20.170	1.800	35.766	1.00 32.90
	ATOM	866	CA	LEU A			21.575	1.513	35.625	1.00 34.69
	ATOM	867	C	LEU A			2.225	2.188	34.422	1.00 35.97
	ATOM	868	ŏ	LEU A			23.434	2.037	34.247	1.00 37.11
35	MOTA	869	CB	LEU A			21.811	0.003	35.568	1.00 34.47
33	ATOM	870	CG	LEU A			21.363	~0.757	36.853	1.00 39.98
		871		LEU A			21.588	-2.261	36.702	1.00 40.37
	MOTA	872		LEU A			22.104	-0.231	38.073	1.00 40.86
	ATOM			THR A			21.460	2.913	33.614	1.00 35.72
40	ATOM	873	N				22.115	3.606	32.468	1.00 34.87
40	ATOM	874	CA	THR A				4.825	33.060	1.00 35.15
	ATOM	875	C	THR A			22.792		34.004	1.00 35.13
	MOTA	876	0	THR A			22.206	5.345		1.00 36.04
	ATOM	877	CB	THR A			21.074	4.013	31.409 30.928	1.00 34.78
45	ATOM	878	OG1	THR A			20.507	2.790	30.928	1.00 34.78
45	ATOM	879	CG2				21.709	4.722		1.00 35.32
	ATOM	880	N	ALA A			23.907	5.292	32.541	
	ATOM	881	CA	ALA A			4.595	6.446	33.150	1.00 36.34 1.00 36.12
•	MOTA	882	С	ALA A			3.662	7.647	33.117	
	ATOM	883	0	ALA A			3.031	7.852	32.074	1.00 35.49
50	ATOM	884	CB	ALA A			5.840	6.710		1.00 36.09
	ATOM	885	N	PRO A			23.556	8.404	34.195	1.00 35.90
	ATOM	886	CA	PRO A			4.273	8.154	35.428	1.00 36.06
	ATOM	887	C,	PRO A			3.617	7.057	36.275	1.00 35.41
	ATOM	888	0	PRO A			22.426	7.141	36.602	1.00 34.13
55	MOTA	889	CB	PRO A			4.167	9.484	36.175	1.00 36.61
	MOTA	890	CG	PRO A			2.880	10.077	35.695	1.00 36.41
	ATOM	891	CD	PRO A	106	2	22.862	9.725	34.223	1.00 36.94
	ATOM	892	N	THR A			4.407	6.045	36.646	1.00 35.50
	ATOM	893	CA	THR A	107		3.833	4.857	37.287	1.00 34.62
60	MOTA	894	С	THR A		2	23.256	5.142	38.651	1.00 34.93
	ATOM	895	0	THR A	107	2	3.639	6.108	39.324	1.00 34.98
	MOTA	896	CB	THR A			4.820	3.666	37.301	1.00 39.51
	ATOM	897	OG1	THR A	107	2	4.114	2.516	37.812	1.00 36.86

	ATOM	898	CG2	THR A	107	26.016	3.957	38.203	1.00 39.22
				GLY A		22.272	4.352	39.091	1.00 35.80
	ATOM	899	N						1.00 35.25
	ATOM	900	CA	GLY A	108	21.709	4.504	40.436	-
	ATOM	901	С	GLY A	108	20.719	5.660	40.527	1.00 35.77
5	ATOM	902	ō	GLY A		20.469	6.140	41.646	1.00 35.22
,			_				6.080	39.427	1.00 34.86
	ATOM	903	N	CYS A		20.099			
	ATOM	904	CA	CYS A	109	19.226	7.257	39.416	1.00 35.48
	ATOM	905	С	CYS A	109	17.997	6.934	38.551	1.00 35.39
				CYS A		18.179	6.397	37.449	1.00 36.31
	ATOM	906	0						1.00 35.03
10	ATOM	907	CB	CYS A	109	19.987	8.395	38.610	
	ATOM	908	SG	CYS A	109	21.181	9.249	39.651	1.00 42.26
	ATOM	909	N	ILE A		16.785	7.158	39.035	1.00 34.76
								38.156	1.00 34.74
	ATOM	910	CA	ILE A		15.613	7.181		
	ATOM	911	С	ILE A	110	15.774	8.456	37.264	1.00 34.48
15	ATOM	912	0	ILE A		16.234	9.494	37.723	1.00 32.60
LJ							7.414		1.00 37.69
	ATOM	913	CB	ILE A		14.315			
	ATOM	914	CG1	ILE A	110	14.117	6.274	39.951	1.00 41.68
	ATOM	915	CG2	ILE A	110	13.100	7.594	38.049	1.00 39.85
				ILE A		13.770	4.943	39.384	1.00 43.84
	ATOM	916							1.00 34.15
20	MOTA	917	N	LYS A		15.373	8.287	36.013	
	ATOM	918	CA	LYS A	111	15.522	9.400	35.086	1.00 35.30
	ATOM		C .	LYS A		14.453	9.355	34.002	1.00 35.93
							8.285	33.702	1.00 35.35
	ATOM	920	0	LYS A		13.942			
	ATOM	921	ÇВ	LYS A	111	16.879	9.488	34.449	1.00 39.11
25	ATOM	922	ÇG	LYS A	111	17.713	8.300	34.171	1.00 44.79
25				LYS A		19,195	8.574	34.494	1.00 41.27
	ATOM	923	CD						1.00 41.21
	ATOM	924	CE	LYS A		19.930	7.270	34.207	
	ATOM	925	NZ	LYS A	111 .	19.943	6.341	35.377	1.00 36.81
	ATOM	926	N	LYS A	112	14.181	10.569	33.482	1.00 36.64
20		-		DIO 1	110	13.194	10.511	32.410	1.00 37.12
30	ATOM	927	CA	LYS A					1.00 38.56
	ATOM	928	С	LYS A	112		11.477	31.283	
	ATOM	929	0	LYS A	112	14.382	12.368	31.442	1.00 37.40
			СВ	LYS A		11.784	10.464	32.883	1.00 45.17
	ATOM	930						34.092	1.00 43.77
	ATOM	931	CG	LYS A		11.274	11.108		
35	ATOM	932	CD	LYS A	112	9.855	10.826	34.592	1.00 42.31
	ATOM	933	CE	LYS A	112	9,625	11.830	35.702	1.00 40.92
						8.251	12.207	36.085	1.00 41.42
	ATOM	934	NZ	LYS A					
	ATOM	935	N	HIS A	113	12.734	11.353	30.256	1.00 38.75
	ATOM	936	CA	HIS A	113	12.848	12.198	29.053	1.00 39.66
40	ATOM	937	С	HIS A		14.221	12.110	28.430	1.00 39.88
70						14.895	13.102	28.197	1.00 40.77
	MOTA	938	0	HIS A					
	MOTA	939	CB	HIS A	113	12.341	13.611	29.366	1.00 38.57
	ATOM	940	CG	HIS A	113	11.036	13.636	30.115	1.00 38.81
		941		HIS A		9.924	12.958	29.655	1.00 42.21
40	ATOM						14.172	31.299	1.00 41.13
45	ATOM	942	CD2	HIS A	113	10.680			
	ATOM	943	CE1	HIS A	113	8.934	13.113	30.518	1.00 37.50
	ATOM	944		HIS A		9.378	13.838	31.529	1.00 43.65
				GLY A		14.678	10.906	28.098	1.00 40.03
	MOTA	945	N					27.465	1.00 42.07
	MOTA	946	CA	GLY A		15.948	10.634	27.405	1.00 42.07
50	MOTA	947	С	GLY A	114	15.982	10.869	25.944	1.00 42.05
-		948	ō	GLY A	114	15.006	10.756	25.196	1.00 43.10
	ATOM						11.210	25.445	1.00 42.78
	ATOM	949	N	TYR A		17.172			
	ATOM	950	CA	TYR A	115	17.369	11.441	24.019	1.00 42.59
	ATOM	951	С	TYR A		18.822	11.155	23.637	1.00 42.83
55				TYR A		19.673	11.182	24.526	1.00 40.37
55	MOTA	952						23.612	1.00 43.99
	MOTA	953	CB	TYR A		17.029	12.863		
	MOTA	954	CG	TYR A	115	17.877	13.912	24.298	1.00 45.80
	ATOM	955		TYR A	115	18.944	14.514	23.648	1.00 47.34
							14.263	25.614	1.00 47.05
	MOTA	956		TYR A		17.628			
60	MOTA	957	CEl	TYR A	115	19.718	15.463	24.291	1.00 47.70
	ATOM	958	CE2	TYR A	115	18.375	15.230	26.260	1.00 47.78
				TYR A		19.398	15.845	25.579	1.00 48.85
	MOTA	959	CZ						1.00 49.73
	MOTA	960	ОН	TYR A	115	20.194	16.754	26.228	1.00 43.73

	MOTA	961	N	THR A		19.071	10.953	22.342	1.00 41.33
	ATOM	962	CA	THR A	116	20.415	10.564	21.938	1.00 42.28
	ATOM	963	С	THR A	116	21.285	11.789	21.697	1.00 42.08
	ATOM	964	0	THR A	116	20.858	12.791	21.136	1.00 42.54
5	MOTA	965	CB	THR A	116	20.365	9.722	20.643	1.00 46.87
•	ATOM	966	OG1	THR A	116	19.589	8.542	20.949	1.00 49.86
	ATOM	967	CG2	THR A	116	21.753	9.272	20.209	1.00 46.71
	ATOM	968	N	VAL A		22.552	11.650	22.045	1.00 40.95
	ATOM	969	CA	VAL A		23.575	12.638	21.686	1.00 40.83
10	ATOM	970	C	VAL A		24.538	11.818	20.830	1.00 41.40
10	MOTA	971	ŏ	VAL A		24.850	10.689	21.222	1.00 41.22
	ATOM	972	CB	VAL A		24.297	13.196	22.928	1.00 39.95
		973		VAL A		25.599	13.915	22.569	1.00 41.09
	ATOM			VAL A		23.375	14.216	23.617	1.00 38.93
15	MOTA	974		GLU A		24.986	12.396	19.713	1.00 41.76
15	MOTA	975	N	GLU A		25.908	11.573	18.905	1.00 42.29
	MOTA	976	CA			27.254	12.265	18.832	1.00 42.60
	ATOM	977	C	GLU A			13.498	18.801	1.00 42.94
	MOTA	978	0_	GLU A		27.288		17.556	1.00 46.34
	MOTA	979	СВ	GLU A		25,266	11.262	16.365	1.00 53.27
20	MOTA	980	CG	GLU A		25.923	11.896		
	ATOM	981	CD	GLU A		25.542	11.270	15.029	1.00 55.37
	ATOM	982		GLU A		24.442	10.707	14.879	1.00 54.40
	ATOM -	983	OE2	GLU A		26.425	11.358	14.155	1.00 57.58
	MOTA	984	N	VAL A		28.322	11.485	18.947	1.00 41.77
25	MOTA	985	CA	VAL A		29.661	12.031	18.909	1.00 42.33
	ATOM	986	C	VAL A	119	30.408	11.363	17.735	1.00 43.93
	ATOM	987	0	VAL A	119	30.499	10.136	17.672	1.00 42.59
	MOTA	988	CB	VAL A	119	30.486	11.790	20.187	1.00 42.91
	MOTA	989	CG1	VAL A	119	31.894	12.330	19.963	1.00 40.51
30	ATOM	990	CG2	VAL A	119	29.868	12.517	21.398	1.00 38.33
	ATOM	991	N	GLN A		30.927	12.212	16.870	1.00 44.65
	ATOM	992	CA	GLN A		31.698	11.727	15.723	1.00 47.49
. •	ATOM	993	C	GLN A		33.182	11,912	16.001	1.00 50.11
	ATOM	994	ō	GLN A		33.676	13.040	16.117	1.00 48.28
35	ATOM	995	CB	GLN A		31.330	12.541	14.481	1.00 48.85
"	ATOM	996	CG	GLN A		29.834	12.547	14.248	1.00 52.37
	ATOM	997	CD	GLN A		29.456	13.089	12.884	1.00 55.34
	MOTA	998		GLN A		30.319	13.536	12.135	1.00 59.08
	MOTA	999	NE2			28.164	13.016	12.580	1.00 56.41
40	ATOM	1000	N	PHE A		33.872	10.771	16.065	1.00 52.87
70		1001	CA	PHE A		35.287	10.824	16.382	1.00 58.51
	MOTA	1001	C	PHE A		36.147	11.092	15.162	1.00 62.63
	MOTA	1002		PHE A		37.367	11.134	15.318	1.00 63.15
	MOTA		O	PHE A		35.764	9.660	17.219	1.00 54.96
15	ATOM	1004	CB	PHE A		35.110	9.596	18.571	1.00 55.06
45	ATOM	1005	CG	PHE A		33.110	8.822	18.763	1.00 54.70
	ATOM	1006				35.627	10.295	19.641	1.00 52.63
	MOTA	1007	CD2	PHE A		33.389	8.726	20.000	1.00 54.52
	MOTA	1008		PHE A			10.209	20.885	1.00 53.27
<b>c</b> 0	ATOM	1009		PHE A		35.032		21.065	
50	ATOM			PHE A	121	33.908			1.00 66.78
	MOTA	1011	N	ASP A	122	35.547	11.249	13.988	1.00 00.70
	ATOM	1012	CA	ASP A		36.246	11.801	12.847	
	ATOM	1013	С	ASP A		35.414	12.152	11.632	1.00 74.59
	ATOM	1014	0	ASP A		35.591	13.228	11.042	1.00 75.06
55	MOTA	1015	CB	ASP A		37.480	10.970	12.495	1.00 80.67
	MOTA	1016	CG	ASP A		38.640	11.910	12.201	1.00 84.98
	MOTA	1017		ASP A		38.366	13.018	11.687	1.00 87.13
	ATOM	1018	OD2	ASP A		39.782	11.512	12.508	1.00 89.25
	ATOM	1019	N	GLY A		34.511	11.277	11.204	1.00 77.06
60	MOTA	1020	CA	GLY A		33.653	11.544	10.050	1.00 79.12
	MOTA	1021	С	GLY A		32.499	10.546	9.975	1.00 80.59
	MOTA	1022	0	GLY A	123	32.702	9.332	10.039	1.00 80.64
	ATOM	1023	N	ASP A	124	31.278	11.063	9.813	1.00 81.67

	MOTA	1024	CA	ASP A	124	30.122	10.180	9.717	1.00 82.77
	ATOM	1025	С	ASP A	124	30.000	9.574	8.324	1.00 82.74
	ATOM	1026	ō	ASP A		29.673	9.777	7.400	0.00 99.00
	MOTA	1027	СВ	ASP A		28.828	10.863	10.141	1.00 87.39
5			CG	ASP A		27.968	9.968	11.019	1.00 91.94
)	MOTA	1028							1.00 93.18
	ATOM	1029		ASP A		28.495	8.933	11.489	
	ATOM	1030	OD2	ASP A		26.779	10.288	11.244	1.00 93.68
	ATOM	1031	N	ASN A	127	34.160	6.401	13.268	1.00 58.22
	MOTA	1032	CA	ASN A	127	33.743	5.964	14.601	1.00 58.81
10	MOTA	1033	С	ASN A	127	32.728	6.983	15.144	1.00 58.26
	ATOM	1034	ō	ASN A		33.132	8.096	15.487	1.00 58.82
	ATOM	1035	СВ	ASN A		34.919	5.955	15.579	1.00 58.33
						35.959	4.889	15.347	1.00 62.63
	MOTA	1036	CG	ASN A					1.00 64.16
	MOTA	1037		ASN A		35.660	3.697	15.490	
15	MOTA	1038	ND2	ASN A	127	37.179	5.307	15.008	1.00 60.87
	ATOM	1039	N	THR A	128	31.462	6.617	15.180	1.00 57.71
	ATOM	1040	CA	THR A	128	30.411	7.522	15.665	1.00 57.23
	ATOM	1041	С	THR A		29.652	6.846	16.795	1.00 56.48
	ATOM	1042	ō	THR A		29.065	5.792	16.534	1.00 57.24
20				THR A		29.452	7.842	14.501	1.00 56.65
20		1043	CB					13.536	1.00 57.99
	ATOM	1044		THR A		30.208	8.579		1.00 56.34
	MOTA	1045	CG2	THR A		28.244	8.653	14.901	
	MOTA	1046	N	MET A		29.705	7.406	18.002	1.00 54.99
	ATOM	1047	CA	MET A	129	29.082	6.817	19.175	1.00 52.19
25	ATOM	1048	С	MET A	129	27.768	7.511	19.533	1.00 50.42
	ATOM	1049	0	MET A		27.588	8.705	19.319	1.00 48.93
	ATOM	1050	CB		129	30.009	6.909	20.391	1.00 58.00
		1051	CG		129	31.164	5.934	20.458	1.00 62.68
	MOTA			MET A		30.652	4.263	20.904	1.00 68.26
20	MOTA	1052	SD					22.510	1.00 68.01
30	ATOM	1053	CE	MET A		29.906	4.518		
	ATOM	1054	N	HIS A		26.859	6.735	20.126	1.00 49.01
	MOTA	1055	.CA	HIS A	130	25.554	7.194	20.571	1.00 47.13
	MOTA	1056	С	HIS A	130	25.485	7.057	22.095	1.00 46.21
	ATOM	1057	0	HIS A	130	26.009	6.118	22.678	1.00 45.61
35	ATOM	1058	CB	HIS A		24.436	6.345	19.956	1.00 53.82
55	MOTA	1059	CG	HIS A		23.967	6.751	18.602	1.00 62.48
		1060		HIS A		24.816	7.119	17.581	1.00 66.97
	MOTA					22.711	6.825	18.084	1.00 65.45
	MOTA	1061		HIS A					1.00 67.80
40	ATOM	1062		HIS A		24.114	7.424	16.505	
40	ATOM	1063	NE2	HIS A		22.831	7.252	16.784	1.00 68.13
	MOTA	1064	N	TYR A	131	24.998	8.097	22.775	1.00 43.95
	MOTA	1065	CA	TYR A	131	24.923	8.130	24.227	1.00 42.51
	ATOM	1066	С	TYR A	131	23.536	8.666	24.576	1.00 41.94
	ATOM	1067	ō	TYR A		22.956	9.497	23.829	1.00 41.77
45	ATOM	1068	СВ	TYR A		25,948	9.151	24.762	1.00 42.89
73		1069	CG	TYR A		27.390	8.830	24.392	1.00 42.97
	MOTA					27.910	9.421	23.240	1.00 42.76
	MOTA	1070	CD1						1.00 43.24
	ATOM	1071	CD2	TYR A		28.205	7.993	25.131	
	ATOM	1072		TYR A		29.207	9.187	22.828	1.00 44.91
50	MOTA	1073	CE2	TYR A		29.502		24.728	
	ATOM	1074	CZ	TYR A	131	29.991	8.365	23.587	1.00 46.72
	ATOM	1075	OH	TYR A		31.295	8.087	23.222	1.00 49.71
	ATOM	1076	N	THR A		22.979	8.297	25.721	1.00 40.16
			CA	THR A		21.666	8.825	26.098	1.00 37.97
55	ATOM	1077					9.967	27.066	1.00 37.49
55	ATOM	1078	Č	THR A	132	21.835			1.00 37.43
	MOTA	1079	0	THR A		22.535	9.779	28.077	
	ATOM	1080	CB	THR A		20.792	7.731	26.746	1.00 38.81
	ATOM	1081	OG1		132	20.603	6.712	25.757	1.00 36.51
	MOTA	1082		THR A	132	19.452	8.267	27.216	1.00 36.18
60	ATOM	1083	N	ASN A	133	21.212	11.104	26.792	1.00 35.61
	ATOM	1084	CA	ASN A		21.188	12.167	27.803	1.00 36.33
			C	ASN A		19.749	12.266	28.321	1.00 35.53
	ATOM	1085				18.880	11.616	27.771	1.00 35.34
	MOTA	1086	0	ASN A	123	10.000	11.010	2 ,	

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	ATOM	1087	CB	ASN A			21.662	13.500	27.280	1.00 36.82
	ATOM	1088	CG	ASN A	133		22.055	14.518	28.327	1.00 42.70
	ATOM	1089	OD1	ASN A	133		22.192	15.696	27.956	1:00 44.62
	ATOM	1090	ND2	ASN A	133		22.275	14.088	29.562	1.00 37.77
5	ATOM	1091	N	TRP A	134		19.546	12.925	29.444	.1.00 35.65
_	ATOM	1092	CA	TRP A			18.238	12.908	30.127	1.00 36.02
	ATOM	1093	C.	TRP A			17.856	14.318	30.510	1.00 36.37
		1094		TRP A			18.686	15.002	31.095	1.00 37.57
	ATOM		0	TRP A			18.363	12.078	31.423	1,00 34.98
10	ATOM	1095	CB				18.900	10.691	31.221	1.00 35.16
10	ATOM	1096	CG	TRP A				10.320	31.363	1.00 36.78
	ATOM	1097		TRP A			20.214		30.845	1.00 37.83
	ATOM	1098	CD2				18.190	9.511		1.00 37.03
	atom	1099	NE1	TRP A			20.350	8.987	31.103	
	ATOM	1100	CE2	TRP A		٠.		8.458	30.802	1.00 34.86
15	ATOM	1101	CE3	TRP A			16.836	9.240	30.593	1.00 35.07
	ATOM	1102	CZ2	TRP A	134		18.770	7.136	30.516	1.00 36.92
	ATOM	1103	CZ3	TRP A	134		16.498	7.932	30.308	1.00 39.28
	ATOM	1104	CH2	TRP A	134		17.449	6.905	30.240	1.00 38.59
	ATOM	1105	N	THR A			16.634	14.807	30.259	1.00 36.82
20	ATOM	1106	CA	THR A			16.349	16.149	30.758	1.00 36.51
20	ATOM	1107	C	THR A			16.094	16.125	32.270	1.00 37.13
	ATOM	1108	ō	THR A			16.234	17.149	32.927	1.00 35.77
	ATOM		СВ	THR A		· .	15.214	16.871	30.031	1.00 41.31
	ATOM	1110	OG1			-	13.979	16.175	30.224	1.00 39.42
25		1111		THR A			15.461	16.910	28.512	1.00 43.30
45	MOTA		N	HIS A			15.650	15.007	32.831	1.00 36.50
	ATOM	1112 1113	CA	HIS A			15.236	14.974	34.227	1.00 37.55
	MOTA			HIS A			16.017	13.856	34.948	1.00 36.41
	ATOM	1114	C O	HIS A			15.783	12.717	34.570	1.00 37.39
30	ATOM	1115		HIS A			13.731	14.716	34.346	1.00 42.28
30	ATOM	1116	CB	HIS A			12.843	15.884	34.044	1.00 45.85
	ATOM	1117	CG				12.928	16.585	32.850	1.00 44.57
	ATOM	1118		HIS A			11.847	16.467	34.751	1.00 47.41
	MOTA	1119		HIS A			12.039	17.564	32.853	1.00 45.33
25	MOTA	1120		HIS A			11.362	17.512	33.987	1.00 49.47
35	MOTA	1121		HIS A				14.223	35.827	1.00 35.44
	MOTA	1122	N	ILE A			16.941		36.526	1.00 34.42
	ATOM	1123	CA	ILE A			17.696	13.165		1.00 34.06
	MOTA	1124	С	ILE A			17.396	13.291	38.023	1.00 34.00
	MOTA	1125	0	ILE A			17.573	14.396	38.537	
40	MOTA	1126	CB	ILE A			19.209	13.314	36.268	1.00 33.02
	ATOM	1127	CG1				19.527	13.053	34.780	1.00 33.98
	ATOM	1128	CG2	ILE A			19.995	12.315	37.126	1.00 34.85
	ATOM	1129	CD1	ILE A			20.948	13.484	34.427	1.00 35.15
	MOTA	1130	N	TYR A	138		16.902	12.217	38.643	1.00 33.90
45	MOTA	1131	CA	TYR A	138		16.497	12.371	40.067	1.00 34.35
	MOTA	1132	C	TYR A	138	٠.	17.618	11.920	40.998	1.00 34.35
	MOTA	1133	0	TYR A	138		17.925	10.740	40.954	1.00 35.52
	ATOM	1134	CB	TYR A	138		15.196	11.604	40.319	1.00 35.05
	ATOM	1135	CG	TYR A	138		14.075	12.281	39.524	1.00 37.41
50	ATOM	1136	CD1	TYR A	138		13.904	11.924	38.203	1.00 38.75
	ATOM	1137		TYR A			13.272	13.267	40.084	1.00 40.04
	ATOM	1138	CE1				12.922	12.525	37.431	1.00 39.64
	ATOM	1139		TYR A			12.281	13.870	39.308	1.00 41.81
	MOTA	1140	CZ	TYR A			12.128	13.492	37.999	1.00 41.16
55		1141	OH	TYR A			11.171	14.050	37.192	1.00 43.58
رر	MOTA	1141	N	ILE A			18.255	12.823	41.721	1.00 34.48
	MOTA		CA	ILE A			19.360	12.458	42.607	1.00 35.01
	MOTA	1143		ILE A			18.756	12.127	43.996	1.00 34.60
	MOTA	1144	C O	ILE A			18.312	13.089	44.613	1.00 35.20
60	MOTA	1145		ILE A			20.353	13.612	42.774	1.00 34.61
UU	MOTA	1146	CB CG1	ILE A			20.926	14.098	41.421	1.00 38.01
	MOTA	1147	CGI	ILE A	130		21.546	13.200	43.637	1.00 37.55
	MOTA	1148					21.340	12.956	40.588	1.00 38.91
	MOTA	1149	CDI	ILE A	123		21.40/	14.770	30.500	2.00 00.71

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	MOTA	1150	N	CYS A			18.581	10.861	44.301		35.26
	MOTA	1151	CA	CYS A			17.938	10.511	45.590		37.11
	MOTA	1152	С	CYS A			18.952	10.174	46.664		37.66
	ATOM	1153	0	CYS A			19.865	9.384	46.448		37.82
5	ATOM	1154	CB	CYS A			17.078	9.251	45.411		42.45
	MOTA	1155	SG	CYS A			15.483	9.689	44.680	1.00	
	MOTA	1156	N	GLU 1			18.799	10.796	47.831		37.14
	ATOM	1157	CA	GLU A			19.664	10.484	48.966		37.88
	MOTA	1158	С	GLU 1			18.845	9.929	50.125		36.63
10	MOTA	1159	0	GLU A			17.678	9.579	49.925		35.72 43.42
	ATOM	1160	CB	GLU A			20.509	11.709	49.297		53.61
	ATOM	1161	CG	GLU A			21.392	12.145	48.129 48.352		58.09
	ATOM	1162	,CD	GLU A		141	22.122	13.446	48.139		57.53
1.0	MOTA	1163		GLU A			21.544	14.528 13.339	48.747	1.00	
15	MOTA	1164		GLU A			23.308	9.883	51.315	1.00	36.78
	MOTA	1165	N	GLU A			19.445 18.783	9.263	52.479		36.84
	ATOM	1166	CA	GLU A			17.515	9.941	52.907		35.85
	MOTA	1167	C	GLU A			16.496	9.267	53.166		36.59
20	ATOM	1168	O	GLU A			19.780	9.360	53.654		43.10
20	ATOM	1169	CB CG	GLU A		142	21.150	8.763	53.466		54.03
	MOTA	1170 1171	CD	GLU A			22.322	9.696	53.311		60.55
	MOTA	1172		GLU A			22.260	10.679	52.543		59.21
	MOTA MOTA	1173		GLU A			23.401	9.470	53.924		64.17
25	ATOM	1174	N	ALA A			17.477	11.272	52.922	1.00	
23	ATOM	1175	CA	ALA A			16.241	11.960	53.308		36.52
	ATOM	1176	C	ALA A			15.739	12.999	52.312		36.98
	ATOM	1177	ŏ	ALA A			15.239	14.019	52.787		38.22
	ATOM	1178	CB	ALA A			16.560	12.620	54.652	1.00	34.45
30	ATOM	1179	N	SER A			16.163	12.881	51.031	1.00	36.01
50	ATOM	1180	CA	SER A			15.754	13.884	50.048	1.00	35.70
	MOTA	1181	C	SER A			15.959	13.477	48.583	1.00	35.68
	MOTA	1182	ō	SER A	<b>A</b> :	144	16.665	12.520	48.259	1.00	34.04
	ATOM	1183	CB	SER A			16.607	15.158	50.246	1.00	39.92
35	ATOM	1184	OG	SER A	<b>A</b> :	144	17.965	14.791	49.965		46.64
	ATOM	1185	N	VAL A	<b>A</b> :	145	15.361	14.287	47.712		36.14
	ATOM	1186	CA	VAL A	Α :	145	15.576	14.054	46.260		35.49
	ATOM	1187	С	VAL A	Α :	145	15.546	15.410	45.563		36.78
	ATOM	1188	0	VAL A	Α :	145	14.806	16.307	45.979	1.00	
40	ATOM	1189	CB	VAL A	4	145	14.580	13.058	45.707		36.75
	ATOM	1190	CG1	VAL A	4	145	13.141	13.446	46.028		36.76
	ATOM	1191	CG2	VAL A			14.730	12.908	44.192	1.00	37.20
	MOTA	1192	N	THR A			16.453	15.595	44.600		37.23
	MOTA	1193	CA	THR A			16.434	16.814	43.769		39.11
45	MOTA	1194	C	THR A			16.529	16.390	42.297		39.36
	MOTA	1195		THR 1			17.361	15.521	41.981		39.50
	MOTA	1196	CB	THR A			17.709	17.657	44.040		43.57 49.29
	ATOM	1197	OG1				17.833	17.886	45.432		46.55
	ATOM	1198	CG2				17.686	18.953	43.244		39.48
50	ATOM	1199	N	VAL A			15.699	16.998	41.454		38.72
	ATOM	1200	CA	VAL A			15.774	16.719	39.433		39.04
	ATOM	1201	C	VAL A			16.800	17.682	39.851		38.28
	MOTA	1202	0	VAL I			16.921	18.842	39.268		40.51
F F	MOTA	1203	CB	VAL A			14.451	16.800 18.209	39.197		42.34
55	MOTA	1204		VAL A			13.871 14.532	16.183	37.873		36.37
	ATOM	1205		VAL A			17.711	17.103	38.634	1.00	39.16
	ATOM	1206	N	VAL A			18.685	17.103	37.910	1.00	38.86
	MOTA	1207	CA	VAL A			18.569	17.615	36.406		38.03
60	ATOM ATOM	1208 1209	0	VAL A			18.093	16.554	35.994	1.00	38.38
UU	ATOM	1210	CB	VAL A			20.117	17.663	38.437	1.00	37.94
	ATOM	1211		VAL 2	٠.	148	20.196	17.921	39.948	1.00	37.81
	ATOM	1212		VAL A			20.543	16.227	38.135		39.08
	71100			1		<del>-</del>					

	ATOM	1213	N	GLU A	149		19.105	18.521	35.575	1.00 37.21
	ATOM	1214	CA	GLU A	149		19.010	18.272	34.118	1.00 37.63
	MOTA	1215	С	GLU A	149		20.309	17.767	33.517	1.00 36.45
	MOTA	1216	٥	GLU A			21.385	18.214	33.912	1.00 37.57
5	ATOM	1217	CB	GLU A			18.519	19.530	33.403	1.00 48.95
	MOTA	1218	CG	GLU A			19.598	20.527	33.046	1.00 53.64
	MOTA	1219	CD	GLU A			19.100	21.544	32.022	1.00 58.00
	ATOM	1220		GLU A			19.440	22.716	32.229	1.00 55.36
	ATOM	1221	QE2				18.363	21.228	31.067	1.00 61.49
10	MOTA	1222	N	GLY A			20.200	16.861	32.559	1.00 37.43
	MOTA	1223	CA	GLY A			21.399	16.289	31.923	1.00 37.28
	MOTA	1224	С	GLY A			21.910	17.392	30.970	1.00 37.55
	ATOM	1225	0	GLY A		•	21.071	18.016	30.338	1.00 37.62
	ATOM	1226	N	GLN A			23.181	17.714	31.014	1.00 37.21 1.00 36.92
15	ATOM	1227	CA	GLN A			23.713	18.812	30.210	-
	ATOM	1228	С	GLN A			24.833	18.247	29.331	1.00 37.46 1.00 35.27
	ATOM	1229	0	GLN A			25.351	17.166	29.610	1.00 37.56
	MOTA	1230	CB	GLN A			24.280	19.937	31.078	1.00 43.46
••	ATOM	1231	CG	GLN A			23.177	20.667	31.852	
20	ATOM	1232	CD	GLN A			23.691	21.877	32.581	1.00 48.78
	MOTA	1233		GLN A			23.987	22.896	31.946	1.00 49.56 1.00 44.12
	MOTA	1234		GLN A			23.820	21.769	33.905	
	ATOM	1235	N	VAL A			25.173	19.028	28.294	1.00 36.05 1.00 36.80
0.5	ATOM	1236	CA	VAL A			26.097	18.488	27.284	1.00 37.21
25	ATOM	1237	C .	VAL A			27.209	19.494	26.986	1.00 37.21
	ATOM	1238	0	VAL A			26.949	20.667	26.795	1.00 30.72
	ATOM	1239	CB	VAL A			25.385	18.266	25.921 25.148	1.00 37.05
	ATOM	1240		VAL A			26.394	17.345	26.088	1.00 42.64
20	MOTA	1241		VAL A			24.181	17.350	27.005	1.00 42.04
30	ATOM	1242	N	ASP A			28.420	19.032 19.530	26.819	1.00 37.01
	ATOM	1243	CA	ASP A			29.712	19.550	25.554	1.00 37.31
	MOTA	1244	C	ASP A			30.452		24.955	1.00 37.68
	MOTA	1245	0	ASP A			30.094	18.059	28.066	1.00 37.00
25	ATOM	1246	CB	ASP A			30.713	19.123 20.514	28.634	1.00 30.75
35	MOTA	1247	CG	ASP A			30.890		27.840	1.00 53.60
	MOTA	1248		ASP A			30.469	21.441	29.695	1.00 41.06
	MOTA	1249		ASP A			31.411	20.767 19.785	25.263	1.00 36.73
	ATOM	1250	N	TYR A			31.550	19.783	24.299	1.00 38.06
40	ATOM	1251	CA	TYR A			32.497	17.950	24.233	1.00 38.77
40	MOTA	1252	C	TYR A			33.107	16.934	24.276	1.00 38.93
	MOTA	1253	0	TYR A			33.386		23.920	1.00 38.36
	ATOM	1254	CB	TYR A			33.624	20.200		1.00 38.72
	ATOM	1255	CG	TYR A			34.637	19.489	23.038 21.714	1.00 39.88
40	MOTA	1256	CD1				34.295	19.193	23.514	1.00 39.51
45	ATOM	1257	CD2	TYR A			35.875	19.104	20.873	1.00 39.95
	ATOM	1258	CE1	TYR A			35.184	18.532	22.687	1.00 41.22
	ATOM	1259	CE2	TYR A			36.759 36.412	18.454 18.174	21.376	1.00 41.17
	MOTA	1260	CZ	TYR A			37.332	17.516	20.610	1.00 42.16
50	MOTA	1261	OH	TYR A			33.319	18.027		
50	ATOM	1262	N	TYR A	155			16.968	27.018	1.00 38.84
	ATOM	1263	CA	TYR A			33.883 32.959	15.819	27.350	1.00 37.80
	MOTA	1264	C	TYR A			33.530	14.735		1.00 37.77
	ATOM	1265	0	TYR A						1.00 40.03
E	ATOM	1266	CB	TYR A			34.500	17.518	28.323 28.031	1.00 43.75
55	ATOM	1267	CG CD1	TYR A			35.364	18.745	28.230	1.00 44.64
	ATOM	1268		TYR A			34.854	20.021 18.604	27.562	1.00 44.86
	MOTA	1269		TYR A			36.655	21.136	27.950	1.00 44.55
	ATOM	1270		TYR A			35.628		27.271	1.00 46.37
40	ATOM	1271		TYR A			37.440 36.920	19.709 20.963	27.485	1.00 48.04
60	MOTA	1272	CZ	TYR A				20.963	27.230	1.00 50.62
	ATOM	1273	OH	TYR A			37.708 31.651	16.005	27.230	1.00 36.89
	ATOM	1274	N	GLY A					27.833	1.00 36.27
	ATOM ,	1275	CA	GLY A	120		30.784	14.894	21.033	1.00 30.27

	ATOM	1276	С	GLY	A	156	29.374	15.297	28.278		36.29
	ATOM	1277	ō	GLY	Α	156	28.845	16.351	27.962	1.00	36.73
	ATOM	1278	N	LEU			28.737	14.367	28.989	1.00	35.47
		1279	CA	LEU			27.410	14.593	29.547	1.00	36.71
~	MOTA			LEU			27.570	14.782	31.061		36.98
5	ATOM	1280	C				28.299	14.009	31.703		37.91
	ATOM	1281	0	LEU					29.348		36.06
	ATOM	1282	CB	LEU			26.472	13.404			39.11
	MOTA	1283	CG	LEU			26.409	12.902	27.878		
	MOTA	1284	CD1	LEU			25.362	11.777	27.835		40.34
10	MOTA	1285	CD2	LEU	Α	157	25.944	14.021	26.948		37.66
	MOTA	1286	N	TYR	Α	158	26.860	15.773	31.583		36.59
	MOTA	1287	CA	TYR	Α	158	27.043	16.018	33.020		37.29
	ATOM	1288	Ċ	TYR			25.778	16.579	33.654		37.06
	ATOM	1289	ō	TYR			24.813	16.941	32.968	1.00	36.78
15	ATOM	1290	СВ	TYR			28.202	17.007	33.172	1.00	37.24
13	MOTA	1291	ÇG	TYR			27.948	18.410	32.664	1.00	38.25
				TYR			27.547	19.427	33.526		37.59
	ATOM	1292					28.158	18.721	31.322		38.51
	MOTA	1293		TYR				20.718	33.056		38.42
	MOTA	1294	CE1	TYR			27.355				38.37
20	MOTA	1295	CE2	TYR			27.955	20.009	30.839		
	MOTA	1296	CZ			158	27.573	21.006	31.711.		38.61
	ATOM	1297	OH	TYR	Α	158	27.359	22.290	31.260		37.90
	ATOM	1298	N	TYR	Α	159	25.799	16.661	34.979		36.65
	MOTA	1299	CA	TYR	Α	159	24.758	17.394	35.694		37.45
25	ATOM	1300	C	TYR			25.493	18.169	36.801		37.16
~-	ATOM	1301	Õ			159	26.659	17.920	37.045		37.16
	ATOM	1302	СВ	-		159	23.638	16.543	36.301	1.00	37.42
		1303	CG			159	24.161	15.441	37.222		37.53
	ATOM						24.429	14.181	36.732		39.31
20	MOTA	1304	CD1				24.352	15.689	38.574		38.15
30	MOTA	1305		TYR				13.169	37.564		39.89
	ATOM	1306	CE1			159	24.902				38.54
	ATOM	1307	CE2			159	24.823	14.699	39.407		39.58
	ATOM	1308	CZ			159	25.101	13.454	38.893		
	ATOM	1309	ОН			159	25.613	12.488	39.741		40.63
35	ATOM	1310	N			160	24.779	19.122	37.356		36.88
	ATOM	1311	CA			160	25.251	19.943	38.452		38.20
	ATOM	1312	C	VAL	Α	160	24.277	19.714	39.629		38.68
	ATOM	1313	0	VAL	A	160	23.078	19.922	39.514		38.89
	MOTA	1314	CB	VAL	Α	160	25.295	21.439	38.094		41.01
40	ATOM	1315	CG1	VAL	Α	160	25.815	22.251	39.288		39.83
	ATOM	1316	CG2	VAL	Α	160	26.254	21.687	36.916		36.61
	ATOM	1317	N			161	24.818	19.208	40.708	1.00	39.08
	ATOM	1318	CA			161	24.018	18.916	41.919	1.00	39.84
	ATOM	1319	C			161	24.734	19.569	43.095	1.00	39.51
45		1320	ŏ			161	25.900	19.316	43.322	1.00	38.92
43	ATOM						23.939	17.418	42.140		37.10
	MOTA	1321	CB			161	23.189	16.976	43.377	1.00	36.87
	ATOM	1322	CG			161		17.363	43.665		42.27
	ATOM	1323		HIS			21.908				38.66
	ATOM	1324		HIS			23.571	16.163	44.374		
50	ATOM	1325		HIS			21.508	16.811	44.805		34.42
	MOTA	1326	NE2	HIS	Α	161	22.503	16.079	45.262		40.76
	ATOM	1327	N	GLU	Α	162	24.031	20.404	43.832		41.45
	ATOM	1328	CA	GLU	Α	162	24.557	21.115	44.998		43.07
	ATOM	1329	С	GLU	Α	162	25.795	21.928	44.616	1.00	42.85
55	ATOM	1330	ō			162	26.806	21.828	45.304	1.00	43.43
23		1331	ČВ			162	24.930	20.138	46.121	1.00	46.97
	ATOM		CG			162	23.750	19.235	46.415		59.55
	ATOM	1332					23.494	18.891	47.854		61.68
	ATOM	1333	CD			162		19.508	48.387		66.02
	MOTA	1334		GLU			22.551				65.49
60	ATOM	1335		GLU			24.226	18.027	48.364		43.79
	ATOM	1336	N			163	25.786	22.518	43.432	1.00	43 61
	ATOM	1337	CA			163	26.930	23.253	42.929	1.00	43.51
	ATOM	1338	C	GLY	Α	163	28.023	22.409	42.306	1.00	43.73

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	MOTA	1339	0	GLY A			28.958	23.011	41.746	1.00 43.41
	ATOM	1340	N	ILE A			27.986	21.079	42.371	1.00 42.17
	MOTA	1341	CA	ILE A			29.078	20.258	41.896	1.00 41.52 1.00 41.11
	MOTA	1342	С	ILE A			28.743	19.687	40.513	1.00 41.11
5	MOTA	1343	0	ILE A			27.677	19.110	40.314	1.00 40.21
	ATOM	1344	CB	ILE A			29.442	19.081	42.820	1.00 46.60
	MOTA	1345	CG1				29.730	19.597	44.228 42.258	1.00 43.25
	ATOM	1346	CG2				30.651	18.346	45.303	1.00 50.89
4.0	ATOM	1347		ILE A			29.708	18.526	39.561	1.00 30.03
10	MOTA	1348	Ŋ	ARG A			29.613	20.004 19.508	38.202	1.00 40.37
	ATOM	1349	CA	ARG A			29.428 29.979	18.082	38.132	1.00 40.18
	MOTA	1350	C	ARG A			31.139	17.889	38.436	1.00 40.87
	MOTA	1351	0	ARG A			30.211	20.389	37.205	1.00 43.98
1.5	MOTA	1352	CB	ARG A			30.190	19.775	35.799	1.00 48.81
15	MOTA	1353	CG	ARG A			31.056	20.614	34.844	1.00 50.67
	ATOM	1354	CD	ARG A			30.374	21.882	34.644	1.00 54.50
	ATOM	1355	NE	ARG A			30.245	22.592	33.535	1.00 51.25
•	ATOM	1356	CZ	ARG A			30.788	22.211	32.399	1.00 52.55
20	ATOM	1357	NUT	ARG A	165		29.552	23.727	33.612	1.00 46.13
20	ATOM	1358					29.159	17.127	37.747	1.00 38.84
	ATOM	1359	N	THR A			29.502	15.722	37.710	1.00 37.91
	MOTA	1360	CA	THR A			29.318	15.152	36.310	1.00 36.66
	MOTA	1361	C	THR F			28.167	15.109	35.857	1.00 36.77
25	MOTA	1362	O	THR F			28.621	14.911	38.700	1.00 42.71
25	MOTA	1363	CB OG1				28.895	15.399	40.034	1.00 43.09
	MOTA	1364 1365	CG2	THR F			28,934	13.427	38.662	1.00 40.93
	MOTA MOTA	1366	N N	TYR F			30.398	14.733	35.667	1.00 36.29
	ATOM		CA	TYR F			30.309	14.112	34.342	1.00 36.95
30		1367 1368	CA	TYR F			29.970	12.639	34.466	1.00 37.76
30	ATOM		õ	TYR A			30.561	11.961	35.335	1.00 39.69
	ATOM	1369 1370	CB	TYR A			31.611	14.231	33.518	1.00 37.91
	ATOM	1371	CG	TYR A			31.797	15.617	32.933	1.00 38.45
	ATOM	1372		TYR A			32.311	16.637	33.726	1.00 39.32
35	MOTA	1373	CD2	TYR F			31.397	15.937	31.646	1.00 39.69
22	MOTA MOTA	1374	CE1	TYR F			32.458	17.919	33.243	1.00 38.98
	ATOM	1375	CE2	TYR F			31.535	17.214	31.133	1.00 39.08
	MOTA	1376	CZ	TYR A			32.064	18.201	31.946	1.00 40.35
	MOTA	1377	OH	TYR A			32.216	19.488	31.494	1.00 39.61
40	MOTA	1378	N	PHE A			28.924	12.172	33.821	1.00 37.39
40	ATOM	1379	CA	PHE A			28.547	10.772	33.818	1.00 38.28
	ATOM	1380	C	PHE A			28.987	10.058	32.559	1.00 39.29
	ATOM	1381	ŏ	PHE A			28.980	8.825	32.494	1.00 38.30
	ATOM	1382	СВ	PHE A		. •	27.085	10.508	34.167	1.00 38.07
45	ATOM	1383	CG	PHE A			26.068	11.226	33.320	1.00 34.93
73	ATOM	1384	CD1				25.596	10.661	32.153	1.00 36.11
	ATOM	1385	CD2	PHE A			25.609	12.470	33.722	1.00 35.04
	ATOM	1386	CE1	PHE A			24.656	11.337	31.364	1.00 34.93
	ATOM	1387	CE2				24.672	13.140	32.951	1.00 35.38
50	ATOM	1388	CZ	PHE A			24.215	12.564	31.799	1.00 30.80
-	ATOM	1389	N	VAL A			29.331	10.849	31.524	1.00 39.59
	ATOM	1390	CA	VAL A			30.019	10.310	30.354	1.00 40.03
	ATOM	1391	C	VAL A	169		31.149	11.322	30.039	1.00 41.32
	ATOM	1392	ō	VAL A			30.904	12.530	30.018	1.00 39.98
55	ATOM	1393	CB	VAL A	169		29.136	10.155	29.112	1.00 39.59
	ATOM	1394	CG1	VAL A			29.988	9.684	27.917	1.00 41.55
	ATOM	1395	CG2	VAL A	169		28.019	9.107	29.242	1.00 38.14
•	ATOM	1396	N	GLN A			32.377	10.844	29.870	1.00 41.92
	ATOM	1397	CA	GLN A			33.464	11.693	29.396	1.00 42.70
60	ATOM	1398	C	GLN A			33.833	11.215	27.990	1.00 43.33
	ATOM	1399	ō	GLN A			34.348	10.097	27.861	1.00 43.22
	ATOM	1400	СВ	GLN A			34.696	11.627	30.293	1.00 39.48
	ATOM	1401	CG	GLN A	170		34.448	12.220	31.676	1.00 45.31

	MOTA	1402	CD	GLN A		35.691		32.542	1.00 46.68
	MOTA	1403		GLN A		35.649		33.717	1.00 52.69
	MOTA	1404	NE2	GLN A	170	36.816		31.998	1.00 47.58
	ATOM	1405	N	PHE A		33.678		26.982	1.00 43.04
5	ATOM	1406	ÇA	PHE A		33.884	1 11.606	25.629	1.00 44.34
	ATOM	1407	С	PHE A	171	35.302		25.340	1.00 45.84
	ATOM	1408	0	PHE A	171	35.508	3 10.343	24.446	1.00 45.10
	ATOM	1409	CB	PHE A	171	33.422	2 12.612	24.597	1.00 39.68
	MOTA	1410	CG	PHE A		31.96	12.978	24.639	1.00 38.24
10	ATOM	1411		PHE A	171	31.016	5 11.986	24.875	1.00 40.47
	ATOM	1412		PHE A		31.530	14.269	24.412	1.00 35.50
	ATOM	1413		PHE A		29.673		24.905	1.00 37.70
	ATOM	1414	CE2	PHE A	171	30.187		24.441	1.00 37.98
	MOTA	1415	CZ	PHE A		29.248		24.690	1.00 37.99
15	ATOM	1416	N	LYS A		36.28		26.080	1.00 45.69
13	ATOM	1417	CA	LYS A		37.660		25.889	1.00 47.54
		1418	C	LYS A		37.949		26.138	1.00 48.27
	MOTA		ŏ	LYS A		38.884		25.570	1.00 48.26
	MOTA	1419		LYS A		38.594		26.708	1.00 52.42
20	ATOM	1420	CB			40.022		26.201	1.00 57.12
20	ATOM	1421	CG	LYS A		40.904		27.157	1.00 64.24
	ATOM	1422	CD	LYS A				27.018	1.00 65.59
	ATOM	1423	CE	LYS A		42.362		26.949	1.00 65.87
	MOTA	1424	NZ	LYS A		43.304		27.009	1.00 48.25
	MOTA	1425	N	ASP A		37.15		-	1.00 49.23
25	MOTA	1426	CA	ASP A		37.212		27.287	
	MOTA	1427	С	ASP A		36.954		26.007	1.00 49.36
	MOTA	1428	0	ASP A		37.788		25.701	1.00 49.13
	MOTA	1429	CB	ASP A		36.26		28.394	1.00 46.27
	ATOM	1430	CG	ASP A		36.62		29.740	1.00 50.82
30	ATOM	1431	ODl	ASP A	173	35.740		30.605	1.00 51.86
	ATOM	1432	OD2	ASP A	173	37.81		29.970	1.00 52.02
	ATOM	1433	N	ASP A	174	35.90	1 7.144	25.248	1.00 49.89
	ATOM	1434	CA	ASP A	174	35.67	6.430	24.007	1.00 50.79
	ATOM	1435	С	ASP A	174	36.64	7 6.784	22.897	1.00 52.00
35	MOTA	1436	0	ASP A	174	37.08	5 5.884	22.165	1.00 51.37
	MOTA	1437	CB	ASP A	174	34.23	6.582	23.539	1.00 51.65
,	ATOM	1438	CG	ASP A		33.27	5.662	24.274	1.00 51.83
	ATOM	1439		ASP A		33.75		24.847	1.00 48.40
	ATOM	1440		ASP A		32.06		24.269	1.00 52.72
40	ATOM	1441	N	ALA A		37.040		22.808	1.00 52.36
10	ATOM	1442	CA	ALA A		38.02		21.821	1.00 53.78
	ATOM	1443	C	ALA A		39.36		21.964	1.00 54.22
	ATOM	1444	Ö	ALA A		39.99		20.974	1.00 54.32
		1445	СВ	ALA A		38.22		21.910	1.00 51.67
45	MOTA	1446	N	GLU A		39.83		23.186	1.00 55.55
43	ATOM		CA	GLU A		41.120	:	23.409	1.00 56.73
	ATOM	1447		GLU A		41.01		23.234	1.00 57.74
	ATOM	1448	C	GLU A		42.02		23.187	1.00 58.14
	ATOM	1449	0			41.65		24.806	1.00 56.85
60	MOTA	1450	CB	GLU A				25.116	1.00 59.30
50	ATOM	1451	CG	GLU A		41.910		26.537	0.00 99.00
	MOTA	1452	CD	GLU A		42.180			0.00 99.00
	MOTA	1453		GLU A		41.740		27.454	0.00 99.00
	MOTA	1454		GLU A		43.07		26.687	
	MOTA	1455	N	LYS A		39.79		23.170	1.00 58.44
55	ATOM	1456	CA	LYS A		39.55		23.024	1.00 59.70
	MOTA	1457	С	LYS A		39.31		21.561	1.00 60.20
	ATOM	1458	0	LYS A	177	39.32		21.198	1.00 60.71
	MOTA	1459	CB	LYS A	177	38.35		23.887	1.00 63.58
	ATOM	1460	CG	LYS A	177	38.05		24.029	1.00 67.87
60	ATOM	1461	CD	LYS A	177	37.04		25.140	1.00 70.02
	ATOM	1462	CE	LYS A	177	36.87		25.242	1.00 73.46
	MOTA	1463	NZ	LYS A	177	36.22	1 -0.638	26.517	1.00 75.57
	ATOM	1464	N	TYR A	178	38.90	8 4.057	20.758	1.00 60.36
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	ATOM	1465	CA	TYR A	178	38.551	3,798	19.381	1.00 61.69
	ATOM	1466	C .	TYR A		39.369	4.563	18.361	1.00 62.23
	ATOM	1467	ō	TYR A		39.821	3.923	17.409	1.00 63.37
	ATOM	1468	CB	TYR A		37.057	3.988	19.100	1.00 61.34
5	MOTA	1469	CG	TYR A		36.139	3.197	20.010	1.00 61.39
-	ATOM	1470	CD1			36.249	1.811	20.081	1.00 62.10
	ATOM	1471	CD2	TYR A	178	35.189	3.821	20.798	1.00 61.28
	MOTA	1472	CE1	TYR A		35.440	1.072	20.929	1.00 62.32
0.2	MOTA	1473	CE2	TYR A		34.378	3.097	21.651	1.00 62.22
10	ATOM	1474	CZ	TYR A		34.505	1.723	21.707	1.00 62.51 1.00 62.15
	ATOM	1475	OH	TYR A		33.696	0.989	22.540 18.525	1.00 62.15
	MOTA	1476	N	SER A		39.573	5.858 6.703	17.498	1.00 64.19
	ATOM	1477	CA	SER A		40.159 41.662	6.923	17.631	1.00 64.98
1.5	ATOM	1478	C	SER A		42.279	6.550	18.630	1.00 64.79
15	MOTA	1479	0	SER A		39.470	8.077	17.505	1.00 65.34
	MOTA	1480	CB	SER A		39.982	8.923	16.491	1.00 69.64
	ATOM	1481	OG N	LYS A		42.254	7.552	16.609	1.00 65.75
	ATOM	1482 1483	CA	LYS A		43.672	7.876	16.658	1.00 66.93
20	ATOM ATOM	1484	C	LYS A		43.886	9.316	17.107	1.00 67.25
20		1485	ŏ	LYS A		44.806	9.591	17.887	1.00 67.85
	ATOM	1486	СВ	LYS A		44.431	7.615	15.360	1.00 69.83
	ATOM	1487	CG	LYS A		45.938	7.673	15.581	1.00 73.65
	ATOM	1488	CD	LYS A		46.681	8.326	14.427	1.00 76.04
25	ATOM	1489	CE	LYS A		48.016	8.889	14.887	1.00 75.99°
	ATOM	1490	NZ	LYS A		47.829	10.056	15.790	1.00 78.28
	ATOM	1491	N	ASN A	181	43.017	10.216	16.652	1.00 66.93
	ATOM	1492	CA.	ASN A	181	43.126	11.609	17.113	1.00 66.52
	ATOM	1493	С	ASN A	181	42.128	11.842	18.240	1.00 65.58
30	MOTA	1494	0	ASN A	181	41.133	11.131	18.348	1.00 65.86
	MOTA	1495	CB	ASN A		43.174	12.454	16.113	0.00 99.00
	ATOM	1496	CG	ASN A		44.528	12.902	15.624	0.00 99.00
	MOTA	1497		ASN A		45.545	12.248	15.833	0.00 99.00 0.00 99.00
	MOTA	1498		ASN A		44.525	14.056	14.935	1.00 65.06
35	ATOM	1499	N	LYS A		42.432	12.756	19.155 20.232	1.00 62.47
	MOTA	1500	CA	LYS A		41.483 40.755	13.086 14.370	19.855	1.00 60.44
	MOTA	1501	C	LYS A		40.755	15.405	20.491	1.00 60.89
	MOTA	1502	0	LYS A		42.257	13.224	21.544	1.00 68.03
40	ATOM	1503	CB	LYS A		41.462	13.626	22.774	1.00 70.69
40	ATOM	1504 1505	CD CD	LYS A		42.223	14.606	23.651	1.00 73.12
	ATOM ATOM	1506	CE	LYS A		41.360	15.246	24.718	1.00 73.28
	ATOM	1507	NZ	LYS A		40.729	16.534	24.325	1.00 74.63
	ATOM	1508	N	VAL A		40.007	14.401	18.761	1.00 58.04
45	MOTA	1509	CA	VAL A		39.240	15.559	18.324	1.00 55.16
	ATOM	1510	C	VAL A	183	37.899	15.073	17.781	1.00 52.42
	ATOM	1511	0	VAL A	183	37.902	14.220	16.898	1.00 52.43
	ATOM	1512	CB	VAL A	183	39.936	16.462	17.302	1.00 58.84
	ATOM	1513	CG1	VAL A	183	38.965	17.219	16.395	1.00 61.12
50	ATOM	1514	CG2	VAL A	183	40.809	17.504	18.010	1.00 60.87
	ATOM	1515	N	TRP A		36.785	15.581	18.302	1.00 49.48
	MOTA	1516	CA	TRP A		35.495	15.042	17.878	1.00 45.59 1.00 44.01
	MOTA	1517	Ç	TRP A	184	34.490	16.159	17.706 17.913	1.00 42.88
~ ~	ATOM	1518	0	TRP A		34.791	17.353	18.839	1.00 42.62
55	MOTA	1519	CB	TRP A	104	35.049	13.926 14.460	20.253	1.00 39.47
	MOTA	1520	CG	TRP A	104	35.075	15.236	20.233	1.00 37.50
	MOTA	1521	CDT	TRP A	104	34.123 36.109	14.271	21.216	1.00 41.84
	ATOM	1522	CD2	TRP A	104	34.491	15.541	22.123	1.00 36.00
60	MOTA	1523		TRP A		35.717	14.963	22.380	1.00 39.37
UU	ATOM ATOM	1524 1525		TRP A		37.328	13.587	21.201	1.00 39.30
	ATOM	1525	C22	TRP A	184	36.504	14.983	23.529	1.00 41.08
	MOTA	1527	C23	TRP A	184	38.105	13.599	22.341	1.00 41.95
	ALVEI,	100,	223						

	MOTA	1528	CH2	TRP	A	184	37.692	14.293	23.486		42.37
	ATOM	1529	N	GLU			33.292	15.809	17.270	1.00	42.34
		1530	CA	GLU .	A	185	32.211	16.770	17.099		42.70
	ATOM	1531	С	GLU	A	185	30.950	16.204	17.765		43.74
5	MOTA	1532	0	GLU	Α	185	30.589	15.062	17.506		43.14
	ATOM	1533 <sup>-</sup>	CB	GLU	A	185	31.903	17.074	15.635		42.03
	ATOM	1534	CG	GLU	A	185	33.023	17.900	14.999	_	38.91
	ATOM	1535	CD	GLU	A	185	32.851	18.134	13.528	_	42.12
	MOTA	1536	OE1	GLU	A	185	31.953	17.527	12.910		40.53
10	MOTA	1537	OE2	GLU			33.641	18.955	13.015		39.53
	ATOM	1538	N	VAL			30.374	17.021	18.653		43.76
	MOTA	1539	CA	VAL			29.211	16.563	19.400		43.51
	MOTA	1540	С	VAL			27.943	17.165	18.791		45.37
	MOTA	1541	٥.	VAL			27.804	18.379	18.647		43.51
15	ATOM	1542	CB	VAL			29.339	17.008	20.872		41.67
	MOTA	1543		VAL			28.159	16.494	21.688		41.68
	ATOM	1544	CG2	VAL			30.697	16.596	21.452		34.89
	MOTA	1545	N	HIS			27.012	16.272	18.454		46.62
	ATOM	1546	CA	HIS			25.679	16.656	18.003		50.05
20	MOTA	1547	C	HIS			24.620	16.257	19.023		52.23
	ATOM	1548	0	HIS			24.447	15.055	19.262		52.60
	MOTA	1549	CB	HIS			25.329	15.949	16.672		51.22 53.27
	MOTA	1550	CG	HIS			26.356	16.159	15.600		55.72
	MOTA	1551		HIS			26.135	16.916	14.467 15.495		54.05
25	ATOM	1552		HIS			27.633	15.710 16.925	13.493		49.24
	ATOM	1553		HIS			27.216		14.329		53.78
	ATOM	1554	NE2	HIS.			28.134	16.205 17.199	19.524	_	54.61
	MOTA	1555	N	ALA			23.836	16.882	20.591		57.68
20	MOTA	1556	CA	ALA			22.883 21.502	17.477	20.393		59.90
30	MOTA	1557	C	ALA			20.747	17.673	21.356		60.86
	MOTA	1558	0	ALA			23.501	17.291	21.929		57.50
	ATOM	1559	CB	ALA			21.107	17.773	19.155		61.42
	ATOM	1560	N	GLY			19.809	18.381	18.867		62.45
35	ATOM	1561	CA	GLY GLY			19.862	19.892	18.710		62.76
33	ATOM	1562	C	GLY			18.863	20.613	18.809		63.40
	ATOM	1563 1564	N O	GLY			21.051	20.429	18.448		62.37
	ATOM	1565	CA	GLY			21.211	21.883	18.338		60.60
	ATOM	1566	CA	GLY			22.551	22.159	17.660		58.62
40	ATOM ATOM	1567	Ö	GLY			23.029	21.274	16.958		58.39
+0	ATOM	1568	N	GLN			23.173	23.285	17.983		56.77
	ATOM	1569	CA	GLN			24.466	23.554	17.336		54.69
	MOTA	1570	C	GLN			25.462	22.451	17.687	1.00	52.57
	ATOM	1571	ō .	GLN			25.599	22.082	18.858	1.00	52.69
45	ATOM	1572	CB	GLN			24.972	24.927	17.760	1.00	58.35
	ATOM	1573	CG	GLN			26.437	25.200	17.502	1.00	62.97
	ATOM	1574	CD	GLN			26.166	26.948	17.561	0.00	99.00
	ATOM	1575	OE1	GLN			25.459	27.683	16.891		99.00
	ATOM	1576	NE2	GLN	A	191	27.070	27.403	18.451	0.00	99.00
50	ATOM	1577	N	VAL .	A	192	26.162	21.943	16.684		49.07
	MOTA	1578	CA	VAL	Α	192	27.264	21.012	16.898		45.59
	ATOM	1579	С	VAL .	A	192	28.237	21.647	17.875		44.47
	MOTA	1580	0	VAL			28.473	22.853	17.803		45.15
	ATOM	1581	CB	VAL	Α	192	27.960	20.717	15.544		42.50
55	MOTA	1582	CG1	VAL	A	192	29.047	19.665	15.684		39.90
	ATOM	1583	CG2	VAL	A	192	26.874	20.237	14.584		40.84
	ATOM	1584	N	ILE	A	193	28.831	20.882	18.775		41.81
	MOTA	1585	CA	ILE	Α	193	29.875	21.406	19.636		39.71
	ATOM	1586	C	ILE	Α	193	31.246	21.009	19.127		40.00
60	ATOM	1587	0	ILE			31.569	19.822	19.009		40.12
	MOTA	1588	CB	ILE			29.715	20.868	21.104		36.14
	ATOM	1589	CG1				28.303	21.218	21.573		38.63
	ATOM	1590	CG2	ILE	A	193	30.765	21.585	21.957	1.00	38.85

					102 -	27 016	20.591	22.922	1.00 41.29
	ATOM	1591		ILE A	193	27.916		18.852	1.00 39.67
	ATOM	1592	N	LEU A		32.075	22.008		1.00 40.23
	MOTA	1593	CA	LEU A		33.436	21.783	18.388	1.00 40.23
	ATOM	1594	С	LEU A		34.455	21.944	19.486	
5	MOTA	1595	0	LEU A		34.136	22.489	20.557	
	ATOM	1596	CB	LEU A		33.741	22.794	17.252	1.00 38.19
	MOTA	1597	CG	LEU A		32.736	22.764	16.101	1.00 42.91
	ATOM	1598	CD1	LEU A	194	33.203	23.702	14.980	1.00 42.33
	ATOM	1599	CD2	LEU A	194	32.593	21.362	15.514	1.00 41.05
10	ATOM	1600	N	CYS A	195	35.658	21.430	19.269	1.00 41.29
	ATOM	1601	CA	CYS A		36.711	21.575	20.273	1.00 43.15
	ATOM	1602	C	CYS A		36.956	23.045	20.562	1.00 45.32
	ATOM	1603	ŏ	CYS A		37.083	23,872	19.671	1.00 44.47
	ATOM	1604	СВ	CYS A		37.981	20.893	19.785	1.00 44.75
15	ATOM	1605	SG	CYS A		39.358	21.057	20.920	1.00 43.19
IJ			N	PRO A		36.918	23.423	21.847	1.00 46.44
	MOTA	1606		PRO A		36.978	24.814	22,245	1.00 47.49
	MOTA	1607	CA			38.376	25.335	22.514	1.00 48.05
	MOTA	1608	C	PRO A			26.531	22.759	1.00 49.91
	MOTA	1609	0	PRO A		38.575		23.515	1.00 48.09
20	MOTA	1610	CB	PRO A		36.123	24.820	24.079	1.00 47.48
	MOTA	1611	CG	PRO A		36.293	23.449		1.00 47.40
	ATOM	1612	CD	PRO A		36.834	22.513	23.018	
	ATOM	1613	N	THR A		39.365	24.477	22.488	1.00 48.20
	ATOM	1614	CA	THR A		40.753	24.821	22.766	1.00 49.51
25	ATOM	1615	С	THR A	197	41.610	24.697	21.491	1.00 48.70
	ATOM	1616	0	THR A	197	41.132	24.143	20.508	1.00 47.51
	ATOM	1617	CB	THR A	197	41.337	23.819	23.789	1.00 54.19
	MOTA	1618	OG1	THR A	197	42.063	22.755	23.133	1.00 60.20
	ATOM	1619	CG2	THR A	197	40.249	23.146	24.620	1.00 60.00
30	MOTA	1620	N	SER A		42.874	25.104	21.600	1.00 48.43
-	MOTA	1621	ÇA	SER A		43.755	25.011	20.443	1.00 49.41
	ATOM	1622	Č	SER A		44.106	23.572	20.088	1.00 51.13
	ATOM	1623	õ	SER A		44.340	22.748	20.974	1.00 50.32
	ATOM	1624	СB	SER A		45.022	25.837	20.600	1.00 41.67
35	ATOM	1625	OG	SER A		44.689	27.176	20.863	1.00 42.21
55		1626	N	VAL A		44.135	23.321	18.783	1.00 51.78
	ATOM		CA	VAL A		44.448	22.018	18.210	1.00 54.21
	ATOM	1627				45.846	22.063	17.593	1.00 55.85
	ATOM	1628	C	VAL A		46.229	23.043	16.958	1.00 55.32
40	MOTA	1629	0	VAL A		43.415	21.673	17.105	1.00 55.83
40	ATOM	1630	CB	VAL A			20.450	16.311	1.00 58.48
	MOTA	1631		VAL A		43.848	21.363	17.794	1.00 58.90
	MOTA	1632		VAL A		42.081			1.00 57.80
	MOTA	1633	N	PHE A		46.643	21.031	17.823	1.00 60.84
	ATOM	1634	CA	PHE A		48.033	20.977	17.458	1.00 63.34
45	ATOM	1635	С.	PHE A		48.501	19.863	16.538	
	MOTA	1636	0	PHE A		47.988	18.771	16.373	1.00 63.71
	ATOM	1637	CB	PHE A		48.976	20.962	18.695	1.00 56.48
	ATOM	1638	CG	PHE A	. 200	49.009	22.350	19.286	1.00 51.93
	ATOM	1639		PHE A		49.867	23.304	18.779	1.00 50.53
50	ATOM	1640	CD2	PHE A	. 200	48.118	22.686	20.298	1.00 52.06
	ATOM	1641	CE1	PHE A	. 200	49.844	24.596	19.289	1.00 48.79
	ATOM	1642	CE2	PHE A	200	48.104	23.972	20.813	1.00 46.67
	ATOM	1643	CZ	PHE A	200	48.952	24.921	20.283	1.00 48.73
	ATOM	1644	N	SER A		49.612	20.201	15.906	1.00 65.54
55	ATOM	1645	CA	SER A		50.520	19.367	15.151	1.00 67.49
~ ~	ATOM	1646	C	SER A		50.543	17.927	15.652	1.00 68.56
				SER A		50.873	17.710	16.841	1.00 69.46
	MOTA	1647	O CB		201	51.928	19.984	15.374	1.00 69.06
	ATOM	1648	CB	SER A		51.799	21.376	15.666	1.00 64.56
60	ATOM	1649	OG			50.201	17.025	14.856	1.00 71.71
60	ATOM	1650	OT	SER A		16.850	8.350	41.749	1.00 33.70
	MOTA	1651		WAT W				36.739	1.00 34.70
	ATOM	1652		WAT W		14.700	3.706	45.725	1.00 35.04
	ATOM	1653	OMO	WAT W	3	23.512	-21.581	73.123	1.00 33.04

						_		16 100	£1 006	1.00 35.32
	ATOM	1654		WAT		4		-16.192	51.826	1.00 35.70
	ATOM	1655		WAT	W	5	27.151	6.021	35.728	1.00 35.70
	ATOM	1656			W	6	35.841	19.641	17.028	1.00 36.41
_	MOTA	1657			W	7	12.924	7.105	31.382	1.00 36.40
5.	ATOM	1,658			W	8	22.396	20.058	35.878	1.00 38.24
	MOTA	1659		WAT		9		-18.568	44.759	1.00 38.55
	ATOM	1660		TAW		10	23.931	7.992	29.610	
	MOTA	1661		WAT		11	32.085	22.981	25.237	1.00 38.76
	MOTA	1662	OMO	TAW	W	12	18.237	6.493	43.333	1.00 38.96
10	MOTA	1663	OWO	WAT	W	13	19.973	21.293	36.711	1.00 38.84
	MOTA	1664	OWO	TAW	W	14	14.757	-3.264	32.978	1.00 39.29
	ATOM	1665	OWO	WAT	W	15	19.948	-2.552	48.084	1.00 39.94
	MOTA	1666	000	TAW	W	16	13.394	-3.627	35.420	1.00 40.05
	ATOM	1667	OWO	WAT	W	17	24.218	1.873	40.503	1.00 40.12
15	ATOM	1668	OWO	WAT	W	18	12.970	-9.846	41.166	1.00 40.24
	ATOM	1669	OWO	WAT	W	19	29.332	23.835	36.728	1.00 40.10
	ATOM	1670	0W0	WAT	W	20	32.982	14.615	37.196	1.00 40.03
	ATOM	1671		WAT	W	21	15.963	-14.411	41.801	1.00 40.85
	ATOM	1672			W	22	36.115	14.124	27.538	1.00 41.62
20	ATOM	1673		WAT		23	36.759	24.316	27.854	1.00 41.58
20	MOTA	1674		WAT		24	24.232	-12.120	40.700	1.00 41.18
	ATOM	1675		WAT		25	20.170	8.808	43.184	1.00 41.55
	ATOM.	1676	-	WAT		26	14.174	8.391	52.525	1.00 43.14
	ATOM	1677		WAT		27	25.412	3.713	30.549	1.00 41.48
25	MOTA	1678		WAT		28	14.723	-7.678	41.271	1.00 41.77
4.5		1679		WAT		29	19.317	13.171	52.356	1.00 41.81
	ATOM	1680		WAT		30	23.266	21.233	27.612	1.00 42.08
	MOTA	1681		WAT		31	11.768	12.568	50.507	1.00 42.04
	ATOM	1682		WAT		32	13.539	-13.131	41.639	1.00 42.18
20	MOTA					33	11.508	10.524	52.379	1.00 42.93
30	MOTA	1683		WAT			3.363	-0.753	38.246	1.00 42.96
	MOTA	1684		WAT		34 35	22.835	-3.224	40.382	1.00 43.01
	ATOM	1685		TAW			26.824	22.825	28.526	1.00 43.00
	MOTA	1686		WAT		36		-1.265	41.392	1.00 43.97
25	ATOM	1687		TAW		37	18.644	-6.966	49.493	1.00 43.69
35	MOTA	1688		WAT		38	8.736		38.985	1.00 43.87
	ATOM	1689		WAT		39	15.651	-14.676 -12.297	37.586	1.00 44.06
	MOTA	1690		WAT		40	8.333		39.161	1.00 44.11
	ATOM	1691		TAW		41	25.939	7.843		1.00 44.27
40	ATOM	1692		WAT		42	10.384	9.683	30.485	1.00 44.34
40	MOTA	1693		TAW		43	0.943	8.083	43.152 43.998	1.00 44.65
	MOTA	1694		TAW		44	21.071	2.692		1.00 44.83
	MOTA	1695		TAW		45	16.203	7.540	48.658	1.00 44.88
	ATOM	1696		WAT		46	21.491	-11.431	40.332	1.00 45.24
	MOTA	1697		WAT		47	21.292	-1.593	41.871	1.00 45.24
45	MOTA	1698		WAT		48	33.051	8.145	30.148	1.00 45.89
	MOTA	1699		TAW		49	11.644	15.303	50.268	
	ATOM	1700		WAT		50	35.864	19.699	14.264	1.00 46.41
	MOTA	1701		WAT		51	7.067	6.695	50.824	1.00 46.58
	ATOM	1702	OM0	WAT	W	52	27.033	16.432	44.207	1.00 47.45
50	MOTA	1703	· OWO	WAT	W	53		-10.902	38.882	1.00 47.44
	MOTA	1704	OWO	WAT	W	54	31.232	24.588	19.293	1.00 47.66
	MOTA	1705	OM0	TAW	W	55	21.781	-3.130	44.031	1.00 47.48
	ATOM	1706	OWO	WAT	W	56	7.169	-27.314	41.711	1.00 48.17
	MOTA	1707	OWO	WAT	W	57	33.861	-1.778	23.374	1.00 48.53
55	ATOM	1708		WAT		58	33.357	8.082	26.362	1.00 48.51
	MOTA	1709		WAT		59	26.396	24.163	13.998	1.00 49.70
	ATOM	1710		WAT		60	21.233	20.044	43.429	1.00 49.73
	ATOM	1711		TAW		61	39.604	24.739	18.318	1.00 49.28
	ATOM	1712		WAT		62	24.974	-18.827	45.601	1.00 50.87
60	MOTA	1713		TAW		63	21.207	-0.654	31.876	1.00 49.81
	ATOM	1714		WAT		64	13.203	8.179	28.792	1.00 50.30
	ATOM	1715		WAT		65	21.887	5.385	43.977	1.00 50.58
	ATOM	1716		TAW		66	24.468	6.206	27.276	1.00 50.23

ATOM 1718 ONO WAT W 68 18.759 17.803 28.596 1.00 50.80 ATOM 1719 ONO WAT W 70 13.821 14.472 25.933 1.00 52.05 1.00 52.05 ATOM 1721 ONO WAT W 71 22.450 0.866 42.331 1.00 52.52 ATOM 1722 ONO WAT W 72 37.480 14.479 34.609 1.00 52.16 ATOM 1723 ONO WAT W 72 37.480 14.4798 12.928 1.00 52.16 ATOM 1723 ONO WAT W 74 33.7480 14.4798 12.928 1.00 52.16 ATOM 1725 ONO WAT W 74 33.7480 14.4798 12.928 1.00 52.80 ATOM 1725 ONO WAT W 74 33.7480 14.4798 34.609 1.00 52.80 ATOM 1725 ONO WAT W 76 8.849 6.339 29.567 1.00 53.08 ATOM 1726 ONO WAT W 77 2.499 1.596 40.507 1.00 53.08 ATOM 1728 ONO WAT W 78 8.453 -4.313 50.928 1.00 53.22 ATOM 1728 ONO WAT W 78 8.453 -4.313 50.928 1.00 53.22 ATOM 1730 ONO WAT W 80 29.311 23.613 25.169 1.00 53.94 ATOM 1731 ONO WAT W 80 29.311 23.613 25.169 1.00 53.94 ATOM 1731 ONO WAT W 80 29.311 23.613 25.169 1.00 53.94 ATOM 1732 ONO WAT W 82 10.533 -28.803 42.420 1.00 54.33 ATOM 1733 ONO WAT W 82 10.533 -28.803 42.420 1.00 54.33 ATOM 1734 ONO WAT W 82 10.533 -28.803 42.420 1.00 54.33 ATOM 1735 ONO WAT W 86 25.591 -14.157 41.467 1.00 54.32 ATOM 1735 ONO WAT W 86 23.172 23.184 42.047 1.00 55.98 ATOM 1735 ONO WAT W 86 23.172 23.184 42.047 1.00 55.98 ATOM 1735 ONO WAT W 88 23.172 23.184 42.047 1.00 55.98 ATOM 1735 ONO WAT W 88 23.172 23.184 42.047 1.00 55.09 ATOM 1738 ONO WAT W 89 29.5591 -14.157 41.467 1.00 54.35 ATOM 1738 ONO WAT W 89 23.05591 4.678 29.129 1.00 54.35 ATOM 1736 ONO WAT W 89 33.899 3.408 32.498 1.00 55.07 ATOM 1736 ONO WAT W 89 33.899 3.408 32.498 1.00 55.07 ATOM 1736 ONO WAT W 99 13.266 -14.306 47.361 1.00 55.07 ATOM 1740 ONO WAT W 99 13.266 -14.306 47.361 1.00 55.07 ATOM 1740 ONO WAT W 99 13.266 -14.306 47.361 1.00 55.07 ATOM 1741 ONO WAT W 99 13.266 -14.306 47.361 1.00 55.07 ATOM 1741 ONO WAT W 99 13.266 -14.306 47.361 1.00 55.07 ATOM 1740 ONO WAT W 99 13.266 -14.306 47.361 1.00 55.07 ATOM 1740 ONO WAT W 99 13.791 -29.887 42.473 1.00 56.88				0110	7.7.R.M	fa.t	67	1	6.159	-6.928	39.274	1.00 50.79
ATOM 1719 ONO NAT W 69 ATOM 1720 ONO NAT W 70 ATOM 1721 ONO NAT W 70 ATOM 1721 ONO NAT W 70 ATOM 1722 ONO NAT W 71 ATOM 1722 ONO NAT W 71 ATOM 1722 ONO NAT W 72 ATOM 1724 ONO NAT W 73 ATOM 1724 ONO NAT W 73 ATOM 1724 ONO NAT W 73 ATOM 1725 ONO NAT W 73 ATOM 1725 ONO NAT W 74 ATOM 1726 ONO NAT W 75 ATOM 1726 ONO NAT W 75 ATOM 1727 ONO NAT W 75 ATOM 1727 ONO NAT W 75 ATOM 1727 ONO NAT W 75 ATOM 1728 ONO NAT W 75 ATOM 1729 ONO NAT W 75 ATOM 1731 ONO NAT W 85 ATOM 1731 ONO NAT W 86 ATOM 1732 ONO NAT W 86 ATOM 1733 ONO NAT W 87 ATOM 1733 ONO NAT W 87 ATOM 1735 ONO NAT W 88 ATOM 1736 ONO NAT W 88 ATOM 1736 ONO NAT W 88 ATOM 1737 ONO NAT W 87 ATOM 1738 ONO NAT W 88 ATOM 1736 ONO NAT W 88 ATOM 1737 ONO NAT W 88 ATOM 1736 ONO NAT W 88 ATOM 1737 ONO NAT W 88 ATOM 1736 ONO NAT W 88 ATOM 1737 ONO NAT W 88 ATOM 1736 ONO NAT W 88 ATOM 1737 ONO NAT W 88 ATOM 1736 ONO NAT W 88 ATOM 1737 ONO NAT W 88 ATOM 1737 ONO NAT W 88 ATOM 1738 ONO NAT W 88 ATOM 1739 ONO NAT W 89 ATOM 1739 ONO NAT W 89 ATOM 1736 ONO NAT W 89 ATOM 1737 ONO NAT W 89 ATOM 1737 ONO NAT W 89 ATOM 1738 ONO NAT W 89 ATOM 1739 ONO NAT W 89 ATOM 1736 ONO NAT W 89 ATOM 1737 ONO NAT W 89 ATOM 1737 ONO NAT W 89 ATOM 1738 ONO NAT W 89 ATOM 1739 ONO NAT W 89												
5 ATOM 1720 ONO WAT W 70 5 992 2 145 50.465 1.00 52.52 ATOM 1721 ONO WAT W 71 22 450 0.866 42.331 1.00 52.52 ATOM 1722 ONO WAT W 72 ATOM 1723 ONO WAT W 73 ATOM 1724 ONO WAT W 73 ATOM 1725 ONO WAT W 73 ATOM 1725 ONO WAT W 73 ATOM 1725 ONO WAT W 74 ATOM 1726 ONO WAT W 75 ATOM 1726 ONO WAT W 76 ATOM 1726 ONO WAT W 76 ATOM 1727 ONO WAT W 76 ATOM 1728 ONO WAT W 77 ATOM 1728 ONO WAT W 78 ATOM 1729 ONO WAT W 78 ATOM 1729 ONO WAT W 78 ATOM 1730 ONO WAT W 78 ATOM 1731 ONO WAT W 80 ATOM 1731 ONO WAT W 81 ATOM 1732 ONO WAT W 82 ATOM 1732 ONO WAT W 82 ATOM 1732 ONO WAT W 82 ATOM 1733 ONO WAT W 82 ATOM 1734 ONO WAT W 84 ATOM 1735 ONO WAT W 84 ATOM 1735 ONO WAT W 84 ATOM 1736 ONO WAT W 84 ATOM 1736 ONO WAT W 86 ATOM 1736 ONO WAT W 86 ATOM 1736 ONO WAT W 86 ATOM 1737 ONO WAT W 86 ATOM 1738 ONO WAT W 86 ATOM 1738 ONO WAT W 86 ATOM 1739 ONO WAT W 87 ATOM 1736 ONO WAT W 86 ATOM 1736 ONO WAT W 87 ATOM 1736 ONO WAT W 86 ATOM 1736 ONO WAT W 87 ATOM 1736 ONO WAT W 88 ATOM 1736 ONO WAT W 87 ATOM 1738 ONO WAT W 88 ATOM 1736 ONO WAT W 88 ATOM 1737 ONO WAT W 88 ATOM 1738 ONO WAT W 88 ATOM 1738 ONO WAT W 89 ATOM 1738 ONO WAT W 88 ATOM 1739 ONO WAT W 88 ATOM 1739 ONO WAT W 88 ATOM 1739 ONO WAT W 89 ATOM 1736 ONO WAT W 89 ATOM 1736 ONO WAT W 89 ATOM 1736 ONO WAT W 89 ATOM 1737 ONO WAT W 89 ATOM 1738 ONO WAT W 89 ATOM 1738 ONO WAT W 89 ATOM 1739 ONO WAT W 89 ATOM 1739 ONO WAT W 89 ATOM 1730 ONO WAT W 89 ATOM												1.00 52.01
S ATOM 1721 OND WAT W 71 22.450 0.866 42.331 1.00 52.86 ATOM 1722 OND WAT W 72 7.480 14.455 29.856 1.00 52.16 ATOM 1724 OWD WAT W 73 7.914 14.799 34.609 1.00 52.16 ATOM 1724 OWD WAT W 74 33.074 14.728 12.928 1.00 52.80 ATOM 1725 OWD WAT W 75 8.849 6.339 29.567 1.00 53.08 ATOM 1726 OWD WAT W 75 8.849 6.339 29.567 1.00 53.08 ATOM 1728 OWD WAT W 77 2.499 9.596 40.507 1.00 53.08 ATOM 1728 OWD WAT W 78 8.453 -4.313 50.928 1.00 53.22 ATOM 1729 OWD WAT W 79 13.988 -16.279 37.477 1.00 54.52 ATOM 1730 OWD WAT W 80 29.311 23.613 25.169 1.00 53.82 ATOM 1730 OWD WAT W 81 10.698 17.607 37.566 1.00 53.87 ATOM 1731 OWD WAT W 82 10.533 -28.803 42.420 1.00 54.31 ATOM 1733 OWD WAT W 83 1.674 -0.659 35.990 1.00 54.31 ATOM 1733 OWD WAT W 85 23.172 23.184 42.047 1.00 53.98 ATOM 1733 OWD WAT W 85 23.172 23.184 42.047 1.00 53.98 ATOM 1733 OWD WAT W 85 23.172 23.184 42.047 1.00 53.98 ATOM 1733 OWD WAT W 85 23.172 23.184 42.047 1.00 53.98 ATOM 1733 OWD WAT W 88 33.071 24.177 22.504 1.00 54.33 ATOM 1733 OWD WAT W 88 33.071 24.177 22.504 1.00 55.39 ATOM 1733 OWD WAT W 88 33.071 24.177 22.504 1.00 55.39 ATOM 1734 OWD WAT W 89 2.865 -1.40.678 27.285 1.00 55.10 ATOM 1734 OWD WAT W 89 2.865 -1.40.678 27.285 1.00 55.07 ATOM 1734 OWD WAT W 99 13.268 -18.762 37.013 1.00 55.07 ATOM 1740 OWD WAT W 99 13.268 -18.762 37.013 1.00 55.07 ATOM 1740 OWD WAT W 99 13.268 -18.762 37.013 1.00 55.07 ATOM 1740 OWD WAT W 99 13.791 -29.887 42.473 1.00 56.24 ATOM 1745 OWD WAT W 99 13.791 -29.887 42.473 1.00 56.35 ATOM 1746 OWD WAT W 99 13.791 -29.887 42.473 1.00 56.35 ATOM 1745 OWD WAT W 99 13.791 -29.887 42.473 1.00 56.35 ATOM 1745 OWD WAT W 99 13.791 -29.887 42.473 1.00 56.35 ATOM 1745 OWD WAT W 99 13.791 -29.887 42.473 1.00 56.35 ATOM 1745 OWD WAT W 99 13.791 -29.887 42.473 1.00 56.35 ATOM 1745 OWD WAT W 101 -0.503 -1.104 43.920 1.00 56.35 ATOM 1745 OWD WAT W 101 -0.503 -1.104 43.920 1.00 56.35 ATOM 1746 OWD WAT W 101 -0.503 -1.104 43.920 1.00 56.35 ATOM 1747 OWD WAT W 101 -0.503 -1.104 43.920 1.00 56.35 ATOM 1747 OWD WAT W 101 -0.503 -1.104 43.920 1.00 56.35												1.00 52.52
ATOM 17122 ONO WAT W 72 ATOM 1723 ONO WAT W 73 ATOM 1724 ONO WAT W 74 ATOM 1724 ONO WAT W 75 ATOM 1725 ONO WAT W 75 ATOM 1725 ONO WAT W 75 ATOM 1726 ONO WAT W 75 ATOM 1726 ONO WAT W 75 ATOM 1727 OWO WAT W 77 ATOM 1729 OWO WAT W 77 ATOM 1729 OWO WAT W 77 ATOM 1729 OWO WAT W 79 ATOM 1730 OWO WAT W 79 ATOM 1730 OWO WAT W 79 ATOM 1730 OWO WAT W 79 ATOM 1731 OWO WAT W 80 ATOM 1731 OWO WAT W 80 ATOM 1731 OWO WAT W 82 ATOM 1733 OWO WAT W 82 ATOM 1733 OWO WAT W 82 ATOM 1733 OWO WAT W 82 ATOM 1734 OWO WAT W 84 ATOM 1735 OWO WAT W 85 ATOM 1735 OWO WAT W 86 ATOM 1736 OWO WAT W 87 ATOM 1736 OWO WAT W 88 ATOM 1736 OWO WAT W 86 ATOM 1736 OWO WAT W 87 ATOM 1736 OWO WAT W 87 ATOM 1736 OWO WAT W 88 ATOM 1736 OWO WAT W 86 ATOM 1736 OWO WAT W 87 ATOM 1736 OWO WAT W 87 ATOM 1736 OWO WAT W 88 ATOM 1736 OWO WAT W 87 ATOM 1736 OWO WAT W 88 ATOM 1736 OWO WAT W 87 ATOM 1736 OWO WAT W 88 ATOM 1736 OWO WAT W 87 ATOM 1736 OWO WAT W 88 ATOM 1736 OWO WAT W 88 ATOM 1736 OWO WAT W 89 ATOM 1738 OWO WAT W 89 ATOM 1740 OWO WAT W 99 ATOM 1740 OWO WAT W 99 ATOM 1740 OWO WAT W 99 ATOM 1744 OWO WAT W 99 ATOM 1745 OWO WAT W 99 ATOM 1746 OWO WAT W 99 ATOM 1746 OWO WAT W 99 ATOM 1747 OWO WAT W 99 ATOM 1747 OWO WAT W 99 ATOM 1748 OWO WAT W 99 ATOM 1748 OWO WAT W 99 ATOM 1749 OWO WAT W 9	5											1.00 52.86
ATOM 17123 OWO WAT W 73 7.914 14.799 34.609 1.00 52.80 ATOM 1724 OWO WAT W 75 -2.139 5.177 35.817 1.00 53.07 ATOM 1726 OWO WAT W 75 -2.139 5.177 35.817 1.00 53.07 ATOM 1728 OWO WAT W 76 8.849 6.339 29.567 1.00 53.08 ATOM 1728 OWO WAT W 77 8.8453 -4.313 50.592 1.00 53.08 ATOM 1728 OWO WAT W 78 8.453 -4.313 50.592 1.00 53.22 ATOM 1729 OWO WAT W 80 13.988 -16.279 37.477 1.00 54.52 ATOM 1730 OWO WAT W 81 10.698 17.607 37.556 1.00 53.94 ATOM 1731 OWO WAT W 81 10.698 17.607 37.556 1.00 53.94 ATOM 1732 OWO WAT W 82 10.533 -28.803 42.420 1.00 54.31 ATOM 1733 OWO WAT W 83 1.674 -0.659 35.990 1.00 54.31 ATOM 1734 OWO WAT W 84 13.238 5.649 29.129 1.00 54.31 ATOM 1735 OWO WAT W 85 23.172 23.184 42.047 1.00 55.98 ATOM 1735 OWO WAT W 86 25.591 -14.157 41.467 1.00 55.39 ATOM 1737 OWO WAT W 87 39.505 4.678 27.285 1.00 55.39 ATOM 1738 OWO WAT W 87 39.505 4.678 27.285 1.00 55.39 ATOM 1737 OWO WAT W 88 33.071 24.177 22.504 1.00 55.39 ATOM 1734 OWO WAT W 89 2.865 -14.306 47.361 1.00 55.09 ATOM 1734 OWO WAT W 89 2.865 -14.306 47.361 1.00 55.09 ATOM 1734 OWO WAT W 89 2.865 -14.306 47.361 1.00 55.09 ATOM 1734 OWO WAT W 89 2.865 -14.306 47.361 1.00 55.09 ATOM 1742 OWO WAT W 99 13.266 -18.762 37.013 1.00 55.07 ATOM 1742 OWO WAT W 99 13.266 -18.762 37.013 1.00 56.24 ATOM 1743 OWO WAT W 99 13.266 -18.762 37.013 1.00 56.24 ATOM 1744 OWO WAT W 99 13.791 -29.887 42.473 1.00 56.35 ATOM 1745 OWO WAT W 99 13.791 -29.887 42.473 1.00 56.35 ATOM 1745 OWO WAT W 99 13.791 -29.887 42.473 1.00 56.35 ATOM 1745 OWO WAT W 99 13.791 -29.887 42.473 1.00 56.35 ATOM 1746 OWO WAT W 101 -0.503 -1.004 43.200 1.00 56.36 ATOM 1749 OWO WAT W 101 -0.503 -1.004 43.200 1.00 56.36 ATOM 1749 OWO WAT W 101 -0.503 -1.004 43.200 1.00 56.36 ATOM 1749 OWO WAT W 101 -0.503 -1.004 43.200 1.00 56.38 ATOM 1749 OWO WAT W 101 -0.503 -1.004 43.200 1.00 56.35 ATOM 1749 OWO WAT W 101 -0.503 -1.004 43.200 1.00 56.35 ATOM 1749 OWO WAT W 101 -0.503 -1.004 43.200 1.00 56.35 ATOM 1749 OWO WAT W 101 -0.503 -1.004 43.200 1.00 56.35 ATOM 1749 OWO WAT W 101 -0.503 -1.004 43.200 1.00 56.35 A	,										29.856	
ATOM 1724 ONO WAT W 74 33.074 14.728 12.928 1.00 52.80 ATOM 1725 ONO WAT W 75 -2.139 5.177 35.817 1.00 53.07 ATOM 1726 ONO WAT W 77 2.499 9.596 40.507 1.00 53.07 ATOM 1729 OWO WAT W 79 13.988 -16.279 37.477 1.00 53.22 ATOM 1729 OWO WAT W 80 29.311 23.613 25.169 1.00 53.24 ATOM 1731 OWO WAT W 81 10.698 17.607 37.556 1.00 53.94 ATOM 1733 OWO WAT W 82 10.533 -28.803 42.420 1.00 54.34 ATOM 1733 OWO WAT W 83 1.674 -0.659 35.990 1.00 54.13 ATOM 1733 OWO WAT W 84 13.238 5.649 29.129 1.00 54.33 ATOM 1735 OWO WAT W 85 23.172 23.184 42.047 1.00 53.94 ATOM 1735 OWO WAT W 86 25.591 -14.157 41.467 1.00 54.33 ATOM 1736 OWO WAT W 88 33.071 24.177 22.855 1.00 55.394 ATOM 1737 OWO WAT W 88 33.071 24.177 22.855 1.00 55.394 ATOM 1738 OWO WAT W 88 33.071 24.177 22.855 1.00 55.104 ATOM 1739 OWO WAT W 89 32.865 -14.306 47.361 1.00 55.093 ATOM 1740 OWO WAT W 89 10.824 5.199 27.990 1.00 54.874 ATOM 1740 OWO WAT W 99 10.824 5.199 27.990 1.00 54.874 ATOM 1740 OWO WAT W 99 13.268 -18.762 37.013 1.00 55.03 ATOM 1740 OWO WAT W 99 13.268 -18.762 37.013 1.00 55.03 ATOM 1740 OWO WAT W 99 13.268 -18.762 37.013 1.00 55.03 ATOM 1744 OWO WAT W 99 13.268 -18.762 37.013 1.00 55.03 ATOM 1746 OWO WAT W 99 13.268 -18.762 37.013 1.00 56.35 ATOM 1746 OWO WAT W 99 13.268 -18.762 37.013 1.00 56.24 ATOM 1746 OWO WAT W 99 13.268 -18.762 37.013 1.00 56.24 ATOM 1746 OWO WAT W 99 13.268 -18.762 37.013 1.00 56.25 ATOM 1746 OWO WAT W 99 13.268 -18.762 37.013 1.00 56.24 ATOM 1746 OWO WAT W 99 13.268 -18.762 37.013 1.00 56.35 ATOM 1746 OWO WAT W 99 13.268 -18.762 37.013 1.00 56.35 ATOM 1746 OWO WAT W 99 13.268 -18.762 37.013 1.00 56.35 ATOM 1746 OWO WAT W 99 13.268 -18.762 37.013 1.00 56.50 ATOM 1746 OWO WAT W 99 13.268 -18.762 37.013 1.00 56.28 ATOM 1746 OWO WAT W 99 13.268 -18.762 37.013 1.00 56.36 ATOM 1746 OWO WAT W 99 13.791 -99.877 1.00 56.35 ATOM 1746 OWO WAT W 99 13.791 -99.877 1.00 56.25 ATOM 1746 OWO WAT W 101 -0.503 -1.004 32.201 1.00 56.55 ATOM 1747 OWO WAT W 101 -0.503 -1.004 32.201 1.00 56.28 ATOM 1746 OWO WAT W 101 -0.503 -1.004 32.201 1.00 56.28 AT											34.609	1.00 52.26
ATOM   1725   ONO WAT W   75   -2.139   5.177   35.817   1.00   53.05   ATOM   1726   ONO WAT W   76   ATOM   1727   ONO WAT W   77   2.499   9.596   40.507   1.00   53.20   ATOM   1728   OWO WAT W   78   8.453   -4.313   50.28   1.00   53.22   ATOM   1730   OWO WAT W   79   13.988   -16.279   37.477   1.00   54.52   ATOM   1731   OWO WAT W   81   10.698   17.607   37.556   1.00   53.24   ATOM   1732   OWO WAT W   82   10.533   -28.803   42.420   1.00   54.32   ATOM   1734   OWO WAT W   83   1.674   -0.659   35.990   1.00   54.11   ATOM   1735   OWO WAT W   85   33.172   23.184   42.047   1.00   54.32   ATOM   1736   OWO WAT W   85   23.172   23.184   42.047   1.00   53.98   ATOM   1737   OWO WAT W   88   33.071   24.177   22.504   1.00   53.98   ATOM   1738   OWO WAT W   88   33.071   24.177   22.504   1.00   53.98   ATOM   1739   OWO WAT W   88   33.071   24.177   22.504   1.00   53.98   ATOM   1739   OWO WAT W   88   33.071   24.177   22.504   1.00   53.98   ATOM   1739   OWO WAT W   89   2.865   -14.306   47.361   1.00   55.09   ATOM   1740   OWO WAT W   90   10.824   5.199   27.990   1.00   54.87   ATOM   1743   OWO WAT W   90   10.824   5.199   27.990   1.00   56.35   ATOM   1744   OWO WAT W   92   19.672   23.135   34.883   1.00   55.09   ATOM   1746   OWO WAT W   93   3.899   3.408   32.498   1.00   56.35   ATOM   1746   OWO WAT W   94   17.326   5.781   50.106   1.00   56.35   ATOM   1746   OWO WAT W   97   16.668   11.173   20.729   1.00   56.28   ATOM   1747   OWO WAT W   98   -0.327   -1.130   37.865   1.00   56.28   ATOM   1750   OWO WAT W   99   3.899   3.408   32.498   1.00   56.28   ATOM   1750   OWO WAT W   99   3.899   3.408   32.498   1.00   56.33   ATOM   1746   OWO WAT W   91   13.266   11.173   20.729   1.00   56.33   ATOM   1746   OWO WAT W   91   13.266   11.173   20.729   1.00   56.35   ATOM   1746   OWO WAT W   91   13.791   -29.887   42.473   1.00   56.58   ATOM   1755   OWO WAT W   99   3.899   3.408   32.498   1.00   56.58   ATOM   1746   OWO WAT W   91   13.791   -29.887   42.473											12.928	
ATOM							75	_	2.139	5.177	35.817	
ATOM 1727 OWO WAT W 77	10 -						76		8.849	6.339	29.567	1.00 53.08
ATOM 1728 OWO WAT W 79 ATOM 1729 OWO WAT W 79 ATOM 1730 OWO WAT W 79 ATOM 1731 OWO WAT W 80 29.311 23.613 25.169 1.00 54.52 ATOM 1732 OWO WAT W 81 10.698 17.607 37.556 1.00 53.87 ATOM 1733 OWO WAT W 82 10.698 17.607 37.556 1.00 53.87 ATOM 1733 OWO WAT W 82 10.698 17.607 37.556 1.00 53.87 ATOM 1733 OWO WAT W 82 10.693 -28.803 42.420 1.00 54.34 ATOM 1734 OWO WAT W 83 1.674 -0.659 35.990 1.00 54.34 ATOM 1735 OWO WAT W 84 13.238 5.649 29.129 1.00 54.35 ATOM 1735 OWO WAT W 85 23.172 23.184 42.047 1.00 53.92 ATOM 1736 OWO WAT W 86 25.591 14.157 41.467 1.00 53.92 ATOM 1738 OWO WAT W 87 ATOM 1738 OWO WAT W 88 33.071 24.177 22.504 1.00 55.92 ATOM 1740 OWO WAT W 89 2.865 -14.306 47.361 1.00 55.09 ATOM 1741 OWO WAT W 91 13.268 -18.762 37.013 1.00 55.07 ATOM 1742 OWO WAT W 92 ATOM 1743 OWO WAT W 93 ATOM 1744 OWO WAT W 93 ATOM 1745 OWO WAT W 93 ATOM 1746 OWO WAT W 94 ATOM 1747 OWO WAT W 95 ATOM 1746 OWO WAT W 95 ATOM 1747 OWO WAT W 96 ATOM 1747 OWO WAT W 97 ATOM 1746 OWO WAT W 99 ATOM 1747 OWO WAT W 99 ATOM 1746 OWO WAT W 99 ATOM 1747 OWO WAT W 99 ATOM 1748 OWO WAT W 99 ATOM 1747 OWO WAT W 99 ATOM 1748 OWO WAT W 99 ATOM 1748 OWO WAT W 99 ATOM 1749 OWO WAT W 98 ATOM 1749 OWO WAT W 98 ATOM 1755 OWO WAT W 102 ATOM 1755 OWO WAT W 102 ATOM 1756 OWO WAT W 103 ATOM 1756 OWO WAT W 104 ATOM 1756 OWO WAT W 104 ATOM 1757 OWO WAT W 107 ATOM 1756 OWO WAT W 109 ATOM 1756 OWO WAT W 109 ATOM 1757 OWO WAT W 109 ATOM 1756 OWO WAT W 119 ATOM 1757 OWO WAT W 119 ATOM 1757 OWO WAT W 119 ATOM 1756 OWO WAT W 119 ATOM 1757 OWO WAT W 120 ATOM 1757 OWO WAT W 121 ATOM 1757 OWO WAT W 122 ATOM 1757 OWO WAT W 123 ATOM 1757 OWO WAT W 124 ATOM 1757 OWO WAT W 126 ATOM 1757 OWO WAT W 127 ATOM 1757 OWO WAT W 128 ATOM 1757 OWO WAT W 128 ATOM 1757 OWO WAT W 129 ATOM 1757 OWO WAT W 129 ATOM 1757 OWO WAT W				OWO	WAT	W	77		2.499	9.596	40.507	
ATOM 1729 OWO WAT W 80 1730 OWO WAT W 80 1731 OWO WAT W 81 10.698 17.607 37.556 1.00 53.94 1736 ATOM 1731 OWO WAT W 82 10.533 -228.803 42.420 1.00 54.31 ATOM 1733 OWO WAT W 83 1.674 -0.659 35.990 1.00 54.31 ATOM 1734 OWO WAT W 84 13.238 5.649 29.129 1.00 54.35 ATOM 1735 OWO WAT W 85 23.172 23.184 42.047 1.00 54.35 ATOM 1736 OWO WAT W 86 25.591 -14.157 41.467 1.00 53.98 ATOM 1737 OWO WAT W 87 ATOM 1738 OWO WAT W 88 ATOM 1738 OWO WAT W 88 ATOM 1738 OWO WAT W 89 ATOM 1738 OWO WAT W 89 ATOM 1739 OWO WAT W 89 ATOM 1730 OWO WAT W 90 ATOM 1740 OWO WAT W 90 10.824 5.199 27.990 1.00 55.07 ATOM 1741 OWO WAT W 91 13.268 -18.762 31.35 34.883 1.00 55.03 ATOM 1743 OWO WAT W 93 ATOM 1743 OWO WAT W 93 ATOM 1743 OWO WAT W 94 ATOM 1744 OWO WAT W 95 ATOM 1745 OWO WAT W 96 ATOM 1746 OWO WAT W 97 ATOM 1747 OWO WAT W 98 ATOM 1746 OWO WAT W 99 ATOM 1746 OWO WAT W 99 ATOM 1746 OWO WAT W 99 ATOM 1747 OWO WAT W 99 ATOM 1746 OWO WAT W 99 ATOM 1747 OWO WAT W 99 ATOM 1748 OWO WAT W 99 ATOM 1748 OWO WAT W 99 ATOM 1749 OWO WAT W 99 ATOM 1746 OWO WAT W 99 ATOM 1747 OWO WAT W 99 ATOM 1747 OWO WAT W 99 ATOM 1748 OWO WAT W 99 ATOM 1748 OWO WAT W 99 ATOM 1749 OWO WAT W 99 ATOM 1749 OWO WAT W 99 ATOM 1750 OWO WAT W 102 ATOM 1750 OWO WAT W 102 ATOM 1751 OWO WAT W 102 ATOM 1752 OWO WAT W 102 ATOM 1753 OWO WAT W 103 ATOM 1754 OWO WAT W 104 ATOM 1755 OWO WAT W 105 ATOM 1756 OWO WAT W 106 ATOM 1757 OWO WAT W 107 ATOM 1758 OWO WAT W 108 ATOM 1758 OWO WAT W 109 ATOM 1759 OWO WAT W 109 ATOM 1759 OWO WAT W 109 ATOM 1750 OWO WAT W 100 ATOM				OWO	WAT	W	78				50.928	
ATOM 1730 OWO WAT W 80 29.311 23.613 25.669 1.00 53.97 ATOM 1731 OWO WAT W 81 10.698 17.607 37.556 1.00 53.97 ATOM 1733 OWO WAT W 82 10.533 -28.803 42.420 1.00 54.31 ATOM 1733 OWO WAT W 83 11.674 -0.659 35.990 1.00 54.31 ATOM 1735 OWO WAT W 85 13.238 5.649 29.129 1.00 54.35 ATOM 1736 OWO WAT W 85 23.172 23.184 42.047 1.00 53.98 ATOM 1737 OWO WAT W 86 25.591 -14.157 41.467 1.00 54.35 ATOM 1737 OWO WAT W 87 39.505 4.678 27.285 1.00 55.10 ATOM 1737 OWO WAT W 88 33.071 24.177 22.504 1.00 53.98 ATOM 1739 OWO WAT W 89 2.865 -14.306 47.361 1.00 55.09 ATOM 1740 OWO WAT W 90 10.824 5.199 27.990 1.00 54.87 ATOM 1740 OWO WAT W 91 13.268 -18.762 37.013 1.00 55.07 ATOM 1742 OWO WAT W 92 19.672 23.135 34.883 1.00 56.03 ATOM 1744 OWO WAT W 94 17.326 5.781 50.106 1.00 56.35 ATOM 1744 OWO WAT W 94 17.326 5.781 50.106 1.00 56.35 ATOM 1747 OWO WAT W 97 16.668 11.173 20.729 1.00 57.28 ATOM 1747 OWO WAT W 99 13.791 20.824 5.199 20.900 1.00 56.83 ATOM 1749 OWO WAT W 99 13.791 20.824 5.199 20.900 1.00 56.35 ATOM 1749 OWO WAT W 99 13.791 20.824 5.199 20.000 56.35 ATOM 1749 OWO WAT W 99 13.791 20.824 5.199 20.000 56.35 ATOM 1749 OWO WAT W 100 14.138 -6.120 36.256 1.00 56.28 ATOM 1750 OWO WAT W 100 14.138 -6.120 36.256 1.00 56.83 ATOM 1750 OWO WAT W 100 14.138 -6.120 36.256 1.00 56.88 ATOM 1755 OWO WAT W 100 39.612 1.004 43.200 1.00 56.88 ATOM 1755 OWO WAT W 100 39.612 1.004 43.200 1.00 57.56 ATOM 1755 OWO WAT W 100 39.612 1.004 43.200 1.00 57.56 ATOM 1755 OWO WAT W 100 25.595 0.503 36.518 1.00 57.76 ATOM 1756 OWO WAT W 103 39.612 16.983 21.891 1.00 57.56 ATOM 1756 OWO WAT W 103 39.612 16.983 21.801 1.00 57.56 ATOM 1756 OWO WAT W 103 39.612 16.983 21.801 1.00 57.56 ATOM 1756 OWO WAT W 103 39.612 16.983 21.801 1.00 58.80 ATOM 1756 OWO WAT W 103 39.612 16.983 21.801 1.00 58.80 ATOM 1756 OWO WAT W 103 39.612 16.983 21.801 1.00 58.80 ATOM 1756 OWO WAT W 103 39.612 16.983 21.201 1.00 58.80 ATOM 1756 OWO WAT W 103 39.612 16.034 12.701 1.00 58.80 ATOM 1756 OWO WAT W 103 39.612 16.034 12.701 1.00 58.80 ATOM 1756 OWO WAT W 103 39.612 16.034 12.7			1729	OWO	WAT	W	79	1	3.988			
ATOM 1732 OWO WAT W 82 10.533 -28.803 42.420 1.00 54.34 ATOM 1733 OWO WAT W 83 1.674 -0.659 35.990 1.00 54.35 ATOM 1735 OWO WAT W 84 13.238 5.649 29.129 1.00 54.35 ATOM 1735 OWO WAT W 85 23.172 23.184 42.047 1.00 53.98 1.00 54.35 ATOM 1736 OWO WAT W 87 39.505 4.678 27.285 1.00 55.10 ATOM 1737 OWO WAT W 88 33.071 24.177 22.504 1.00 53.98 ATOM 1739 OWO WAT W 89 2.865 -14.357 41.467 1.00 53.98 ATOM 1739 OWO WAT W 89 2.865 -14.306 47.361 1.00 55.09 ATOM 1740 OWO WAT W 90 10.824 5.199 27.990 1.00 54.87 ATOM 1740 OWO WAT W 91 13.268 -18.762 37.013 1.00 55.09 ATOM 1740 OWO WAT W 92 19.672 23.135 34.883 1.00 56.03 ATOM 1740 OWO WAT W 93 3.899 3.408 22.498 1.00 56.03 ATOM 1740 OWO WAT W 94 17.326 5.781 50.106 1.00 56.35 ATOM 1745 OWO WAT W 95 46.420 28.753 20.001 1.00 56.50 ATOM 1745 OWO WAT W 99 16.668 11.173 20.729 1.00 57.28 ATOM 1749 OWO WAT W 99 13.91 -29.887 42.473 1.00 56.28 ATOM 1749 OWO WAT W 99 13.791 -29.887 42.473 1.00 56.28 ATOM 1749 OWO WAT W 99 13.791 -29.887 42.473 1.00 56.78 ATOM 1750 OWO WAT W 100 14.138 -6.120 36.256 1.00 56.78 ATOM 1751 OWO WAT W 100 14.138 -6.120 36.256 1.00 56.78 ATOM 1753 OWO WAT W 100 14.138 -6.120 36.256 1.00 56.78 ATOM 1755 OWO WAT W 100 14.138 -6.120 36.256 1.00 56.78 ATOM 1755 OWO WAT W 100 14.138 -6.120 36.256 1.00 56.78 ATOM 1755 OWO WAT W 100 14.138 -6.120 36.256 1.00 56.78 ATOM 1755 OWO WAT W 100 27.034 2.268 29.920 1.00 57.78 ATOM 1755 OWO WAT W 100 27.034 2.268 29.920 1.00 57.78 ATOM 1755 OWO WAT W 100 25.505 0.503 36.518 1.00 56.78 ATOM 1755 OWO WAT W 100 25.505 0.503 36.518 1.00 56.78 ATOM 1756 OWO WAT W 105 21.186 21.778 39.197 1.00 58.80 ATOM 1757 OWO WAT W 105 21.186 21.778 39.197 1.00 58.80 ATOM 1756 OWO WAT W 105 21.186 21.778 39.197 1.00 58.80 ATOM 1756 OWO WAT W 105 21.186 21.778 39.197 1.00 58.80 ATOM 1757 OWO WAT W 105 21.186 21.778 39.197 1.00 58.80 ATOM 1750 OWO WAT W 105 21.186 21.778 39.197 1.00 58.80 ATOM 1750 OWO WAT W 105 21.186 21.778 39.197 1.00 58.80 ATOM 1750 OWO WAT W 105 21.186 21.778 39.197 1.00 58.80 ATOM 1750 OWO WAT W 105 21.866 0.374 1.279 1			1730	OWO	WAT	W	80					
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ATOM 1734 OWO WAT W 84 13.238 5.649 29.129 1.00 54.35 ATOM 1736 OWO WAT W 85 23.172 23.184 42.047 1.00 53.98 ATOM 1736 OWO WAT W 86 25.591 -14.157 41.467 1.00 55.99 ATOM 1738 OWO WAT W 87 39.505 4.678 27.285 1.00 55.10 ATOM 1738 OWO WAT W 89 2.865 -14.306 47.361 1.00 55.99 ATOM 1740 OWO WAT W 99 10.824 5.199 27.990 1.00 55.07 ATOM 1741 OWO WAT W 99 10.824 5.199 27.990 1.00 56.24 ATOM 1743 OWO WAT W 99 13.268 -18.762 37.013 1.00 55.07 ATOM 1744 OWO WAT W 99 13.68 -18.762 37.013 1.00 56.24 ATOM 1745 OWO WAT W 99 4 17.326 5.781 50.106 1.00 56.35 ATOM 1746 OWO WAT W 95 46.420 28.753 20.001 1.00 56.54 ATOM 1747 OWO WAT W 97 16.668 11.173 20.729 1.00 56.58 ATOM 1747 OWO WAT W 97 16.668 11.173 20.729 1.00 56.28 ATOM 1749 OWO WAT W 99 13.791 -29.887 42.473 1.00 56.38 ATOM 1749 OWO WAT W 99 13.791 -29.887 42.473 1.00 56.28 ATOM 1749 OWO WAT W 99 13.791 -29.887 42.473 1.00 56.28 ATOM 1750 OWO WAT W 101 1.039 13.791 -29.887 42.473 1.00 56.88 ATOM 1750 OWO WAT W 101 1.039 13.791 -29.887 42.473 1.00 56.88 ATOM 1753 OWO WAT W 102 7.034 2.268 29.920 1.00 57.28 ATOM 1755 OWO WAT W 103 39.612 16.983 21.840 1.00 57.56 ATOM 1755 OWO WAT W 104 12.322 -7.683 34.772 1.00 56.88 ATOM 1755 OWO WAT W 105 23.186 21.778 39.197 1.00 58.04 ATOM 1755 OWO WAT W 106 25.213 27.127 14.493 1.00 58.04 ATOM 1755 OWO WAT W 106 25.213 27.127 14.493 1.00 58.04 ATOM 1755 OWO WAT W 106 25.213 27.127 14.493 1.00 58.04 ATOM 1755 OWO WAT W 106 25.213 27.127 14.493 1.00 58.04 ATOM 1750 OWO WAT W 106 25.213 27.127 14.493 1.00 58.98 ATOM 1760 OWO WAT W 107 37.899 17.600 47.916 1.00 58.92 ATOM 1760 OWO WAT W 111 23.922 24.315 1.00 58.91 ATOM 1760 OWO WAT W 112 30.025 16.034 17.70 17.00 WAT W 116 23.922 24.315 1.00 58.92 ATOM 1760 OWO WAT W 118 23.932 23.123 29.269 1.00 58.80 ATOM 1760 OWO WAT W 118 23.932 23.123 29.269 1.00 58.80 ATOM 1760 OWO WAT W 118 23.932 23.123 29.269 1.00 58.80 ATOM 1760 OWO WAT W 118 23.932 23.123 29.269 1.00 58.92 ATOM 1763 OWO WAT W 118 23.932 23.123 29.269 1.00 58.93 ATOM 1760 OWO WAT W 118 23.932 23.123 29.269 1.00 58.93 ATOM 1760		ATOM	1732	OWO	TAW	W	82					
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		ATOM	1779					1	3.030	-13.947	38.339	1.00 62.52

					,							
	MOTA	1780	OWO	TAW	W	130	9.7	47	-0.992	36.212	_	59.32
	MOTA	1781	OWO		W	131	24.83	14 -	-11.997	45.661		62.19
	ATOM	1782			W	132	23.20	00	4.574	23.546	1.00	61.90
	ATOM	1783	OWO	WAT	W	133	24.93	38	30.370	17.496		62.23
5	ATOM	1784		TAW	W	134	35.4	59	1.260	16.603	1.00	62.66
•	ATOM	1785	OWO			135	24.1	78	20.068	20.090	1.00	61.73
	ATOM	1786		WAT		136	40.13	27	0.350	18.771	1.00	62.44
	ATOM	1787		WAT		137	19.2	79	14.663	46.778	1.00	63.59
	ATOM	1788		WAT		138	20.0		20.354	46.023	1.00	62.81
10	ATOM	1789		WAT			15.2		18.974	46.516	1.00	63.68
10	MOTA	1790				140	21.2		-25.030	39.386	1.00	63.31
	ATOM	1791				141	26.1		2.756	33.033	1.00	63.89
	ATOM	1792				142	13.2		16.259	48.398	1.00	64.51
	ATOM	1793	OWO			143	23.4		19.596	51.112	1.00	
15	ATOM	1794				144	6.7		10.141	28.981	1.00	
15		1795			W	145	23.6		15.848	49.685		64.50
	ATOM	1796				146	21.8		-6.518	36.556		65.24
	ATOM			WAT		147	10.1		-10.444	36.806	1.00	
	ATOM	1797				148	32.4		10.232	33.677		64.60
20	ATOM	1798		WAT			31.6		6.263	27.510		64.29
20	MOTA	1799		WAT		149			-20.934	39.031		66.94
	ATOM	1800		WAT			4.5			18.085		65.84
	MOTA	1801		WAT		151	38.4		11.674		1.00	
	ATOM	1802				152	42.4		8.992	21.219	1.00	
	MOTA	1803				153	33.9		24.173	27.259		
25	MOTA	1804		WAT		154	24.5		-9.268	45.286	1.00	
	MOTA	1805				155	-2.1	_	-26.008	50.039		66.08
	MOTA	1806		WAT		156	9.0		-32.481	39.896		66.14
	ATOM	1807				157	-3.2		-18.835	45.004	1.00	
	MOTA	1808				158	-3.3		4.020	40.260	1.00	
30	MOTA	1809				159	25.8		24.231	20.622		68.10
	ATOM	1810		TAW			27.1		4.546	24.805		67.75
	MOTA	1811				161	24.0		24.303	35.784	1.00	
	ATOM	1812	OWO	WAT	W	162	7.7		17.585	52.663		70.19
	ATOM	1813	OWO	WAT	W	163	19.3		4.980	47.873	1.00	
35	ATOM	1814	OWO	WAT	W	164	10.43		-4.135	32.539		65.45
	MOTA	1815		WAT	W	165	23.7	98	-0.930	41.113		68.64
	MOTA	1816	OWO	WAT	W	166	2.4		5.318	30.549	1.00	
	ATOM	1817			W	167	9.6	65 ·	-14.876	35.700		65.21
	MOTA	1818	OWO	WAT	W	168	1.7	59	10.431	44.227		69.25
40	MOTA	1819	OWO	WAT	W	169	20.9	60	4.214	26.258		69.97
	ATOM	1820	OWO	TAW	W	170	28.7	69	24.807	27.878	1.00	67.86
	ATOM	1821				171	30.2	12	14.473	8.293	1.00	69.23
	ATOM	1822	OWO	WAT	W	172	20.1	78	0.312	50.589	1.00	70.29
	ATOM	1823	OWO	WAT	W	173	. 19.7	36	6.852	23.117	1.00	70.72
45	ATOM	1824		WAT		174	8.9		16.807	50.514	1.00	70.10
	ATOM	1825				175	25.1		-1.759	34.429	1.00	71.96
	MOTA	1826				176	26.9		25.298	35.563	1.00	68.69
	ATOM	1827	-		W	177	44.9		5.619	13.054	1.00	70.16
	ATOM	1828				178	22.3		24.094	38.170		71.66
50	ATOM	1829	_	WAT		179	-0.6		10.187	33.201	1.00	72.23
J-0	MOTA	1830		WAT		180	11.0		17.856	47.520	1.00	71.42
	MOTA	1831		WAT		181	7.7		0.898	52.950		71.64
	ATOM	1832		WAT					-28.368	52.511		70.10
	AION	1026	240	****	••	~~~	J. 4			<del></del>		

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